(19) World Intellectual Property Organization International Bureau





(43) International Publication Date 10 January 2002 (10.01.2002)

(10) International Publication Number WO 02/02641 A1

(51) International Patent Classification?:	C07K 16/00	Cambridge Antibody Technology Group plc, The Science Park, Melbourn, Nr Royston, Cambridgeshire SG8 6JJ
(21) International Application Number:	PCT/US01/19110	(GB). HILBERT, David [US/US]; 8501 Meaowlark Lane, Bethesda, MD 20817 (US).
(22) International Filing Date: 15 June	2001 (15.06.2001)	(TA) A WOOMED II A TO THE TOTAL THE TOTAL TO THE TOTAL TOTAL TO THE TO
(25) Filing Language:	English	(74) Agents: HOOVER, Kenley, K. et al.; Human Genome Sciences, Inc., 9410 Key West Avenue, Rockville, MD 20850 (US).

(30)	Priority Data:		
	60/212,210	16 June 2000 (16.06.2000)	US
	60/240,816	17 October 2000 (17.10.2000)	US
	60/276,248	16 March 2001 (16.03.2001)	US
	60/277,379	21 March 2001 (21.03.2001)	US
	60/293,499	25 May 2001 (25.05.2001)	US

AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA,

(81) Designated States (national): AE, AG, AL, AM, AT, AU,

- (71) Applicants (for all designated States except US): HUMAN GENOME SCIENCES, INC. [US/US]; 9410 Key West Avenue, Rockville, MD 20850 (US). CAMBRIDGE AN-TIBODY TECHNOLOGY GROUP PLC [GB/GB]; The Science Park, Melbourn, Nr Royston, Cambridgeshire SG8. 6JJ (GB).
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

(72) Inventors; and

(26) Publication Language:

Published:

(75) Inventors/Applicants (for US only): RUBEN, Steven. M. [US/US]; 18528 Heritage Hills Drive, Olney, MD 20832 (US). BARASH, Steven, C. [US/US]; 111 Watkins Pond Blvd., #301, Rockville, MD 20850 (US). CHOI, Gil, H. [KR/US]; 11429 Potomac Oaks Drive, Rockville, MD 20850 (US). VAUGHAN, Tristan [GB/GB]; c/o

with international search report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: ANTIBODIES THAT IMMUNOSPECIFICALLY BIND TO BLYS

(57) Abstract: The present invention relates to antibodies and related molecules that immunospecifically bind to BLyS. The present invention also relates to methods and compositions for detecting or diagnosing a disease or disorder associated with aberrant BLyS expression or inappropriate function of BLyS comprising antibodies or fragments or variants thereof or related molecules that immunospecifically bind to BLyS. The present invention further relates to methods and compositions for preventing, treating or ameliorating a disease or disorder associated with aberrant BLyS expression or inappropriate BLyS function comprising administering to an animal an effective amount of oune or more antibodies or fragments or variants thereof or related molecules that immunospecifically bind to BLyS.



ANTIBODIES THAT IMMUNOSPECIFICALLY BIND TO BLyS

INTRODUCTION

[001] The present invention relates to antibodies and related molecules that immunospecifically bind to BLyS. The present invention also relates to methods and compositions for detecting, diagnosing, or prognosing a disease or disorder associated with aberrant BLyS or BLyS receptor expression or inappropriate function of BLyS or BLyS receptor, comprising antibodies or fragments or variants thereof, or related molecules, that immunospecifically bind to BLyS. The present invention further relates to methods and compositions for preventing, treating or ameliorating a disease or disorder associated with aberrant BLyS or BLyS receptor expression or inappropriate BLyS function or BLyS receptor function, comprising administering to an animal, preferably a human, an effective amount of one or more antibodies or fragments or variants thereof, or related molecules, that immunospecifically bind to BLyS.

BACKGROUND OF THE INVENTION

[002] B lymphocyte stimulator (BLyS) is a member of the tumor necrosis factor ("TNF") superfamily that induces both *in vivo* and *in vitro* B cell proliferation and differentiation (Moore *et al.*, Science 285: 260-263 (1999)). BLyS is distinguishable from other B cell growth and differentiation factors such as IL-2, IL-4, IL-5, IL-6, IL-7, IL-13, IL-15, CD40L, or CD27L (CD70) by its monocyte-specific gene and protein expression pattern and its specific receptor distribution and biological activity on B lymphocytes. BLyS expression is not detected on natural killer ("NK") cells, T cells or B cells, but is restricted to cells of myeloid origin. BLyS expression on resting monocytes is upregulated by interferon-gamma (IFN-gamma). The gene encoding BLyS has been mapped to chromosome 13q34.

[003] BLyS is expressed as a 285 amino acid type II membrane-bound polypeptide and a soluble 152 amino acid polypeptide (Moore *et al.*, 1999 *supra*). The membrane-bound form of BLyS has a predicted transmembrane spanning domain between amino acid residues 47 and 73. The NH₂-terminus of the soluble form of BLyS begins at Ala¹³⁴ of the membrane-bound form of BLyS. Soluble recombinant BLyS has

been shown to induce *in vitro* proliferation of murine splenic B cells and to bind to a cell-surface receptor on these cells (Moore *et al.*, 1999 *supra*). Soluble BLyS administration to mice has been shown to result in an increase in the proportion of CD45R^{dull}, Ly6D^{bright} (also known as ThB) B cells and an increase in serum IgM and IgA levels (Moore *et al.*, 1999 *supra*). Thus, BLyS displays a B cell tropism in both its receptor distribution and biological activity.

Based upon its expression pattern and biological activity, BLyS has been suggested to be involved in the exchange of signals between B cells and monocytes or their differentiated progeny. The restricted expression patterns of BLyS receptor and ligand suggest that BLyS may function as a regulator of T cell-independent responses in a manner analogous to that of CD40 and CD40L in T cell-dependent antigen activation. As such, antibodies and related molecules that immunospecifically bind to BLyS may find medical utility in, for example, the treatment of B cell disorders associated with autoimmunity, neoplasia, or immunodeficiency syndromes.

SUMMARY OF THE INVENTION

The present invention encompasses antibodies (including molecules [005]comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or polypeptide fragment of BLyS. In particular, the invention encompasses antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or polypeptide fragment of human BLyS (SEQ ID NOS:3228 and/or 3229) or BLyS expressed on human monocytes; murine BLyS (SEQ ID NOS:3230 and/or 3231) or BLyS expressed on murine monocytes; rat BLyS (either the soluble forms as given in SEQ ID NOS:3232, 3233, 3234 and/or 3235 or in a membrane associated form, e.g., on the surface of rat monocytes); or monkey BLyS (e.g., the monkey BLyS polypeptides of SEQ ID NOS:3236 and/or 3237, the soluble form of monkey BLyS, or BLyS expressed on monkey monocytes), preferably human BLyS. The present invention also encompasses methods and compositions for detecting, diagnosing, or prognosing diseases or disorders associated with aberrant BLyS or BLyS receptor expression or inappropriate function of BLyS or BLyS receptor in an animal, preferably a mammal, and most preferably a human,

comprising, or alternatively consisting of, use of antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS. Diseases and disorders which can be detected. diagnosed, or prognosed with the antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) of the invention include, but are not limited to, immune disorders (e.g., lupus, rheumatoid arthritis, multiple sclerosis, myasthenia gravis, Hashimoto's disease, and immunodeficiency syndrome), inflammatory disorders (e.g., asthma, allergic disorders, and rheumatoid arthritis), infectious diseases (e.g., AIDS), and proliferative disorders (e.g., leukemia, carcinoma, and lymphoma). The present invention further encompasses methods and compositions for preventing, treating or ameliorating diseases or disorders associated with aberrant BLyS or BLyS receptor expression or inappropriate function of BLyS or BLyS receptor in an animal, preferably a mammal, and most preferably a human, comprising, or alternatively consisting of, administering to said animal an effective amount of one or more antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS. Diseases and disorders which can be prevented, treated or ameliorated by administering an effective amount of an antibody of the invention include, but are not limited to, immune disorders (e.g., lupus, rheumatoid arthritis, multiple sclerosis, myasthenia gravis, Hashimoto's disease, and immunodeficiency syndrome), inflammatory disorders (e.g., asthma, allergic disorders, and rheumatoid arthritis), infectious diseases (e.g., AIDS), and proliferative disorders (e.g., leukemia, carcinoma, and lymphoma).

[006] Using phage display technology, the present inventors have identified single chain antibody molecules ("scFvs") that immunospecifically bind to BLyS, including scFvs that immunospecifically bind to soluble BLyS, scFvs that immunospecifically bind the membrane-bound form of BLyS, and scFvs that immunospecifically bind to both the soluble form and the membrane-bound form of BLyS. Molecules comprising, or alternatively consisting of, fragments or variants of these scFvs (e.g., including VH domains, VH CDRs, VL domains, or VL CDRs having an amino acid sequence of any one of those referred to in Table 1), that immunospecifically bind the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the soluble form and membrane-bound form of BLyS, are also encompassed by the invention,

as are nucleic acid molecules that encode these scFvs, and/or molecules.

In particular, the invention relates to scFvs comprising, or alternatively [007] consisting of, an amino acid sequence selected from the group consisting of SEQ ID NOS: 1-2128, preferably SEQ ID NOS:834 - 872, 1570 - 1595, and 1886 - 1908, and most preferably SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, and 1881 - 1885, as referred to in Table 1 below. In specific embodiments, the present invention relates to scFvs that immunospecifically bind the soluble form of BLyS, said scFvs comprising, or alternatively consisting of, an amino acid sequence of SEQ ID NOS: 1563 - 1569. preferably SEQ ID NOS:1570 - 1595, and most preferably SEQ ID NOS: 1563 - 1569, as referred to in Table 1, below. In other embodiments, the present invention also relates to scFvs that immunospecifically bind the membrane-bound form of BLyS, said scFvs comprising, or alternatively consisting of, an amino acid sequence of SEQ ID NOS: 1881 -2128, preferably SEO ID NOS:1886 - 1908, and most preferably SEO ID NOS: 1881 -1885, as referred to in Table 1 below. The present invention further relates to scFvs that immunospecifically bind both the membrane-bound form and soluble form of BLyS, said scFvs comprising, or alternatively consisting of, an amino acid sequence of SEO ID NOS: 1 - 1562, preferably SEQ ID NOS: 834 - 872, and most preferably SEQ ID NOS: 1 - 46, and 321 - 329, as referred to in Table 1 below. Molecules comprising, or alternatively consisting of, fragments or variants of these scFvs (e.g., including VH domains, VH CDRs, VL domains, or VL CDRs having an amino acid sequence of any one of those referred to in Table 1), that immunospecifically bind the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the soluble form and membrane-bound form of BLyS, are also encompassed by the invention, as are nucleic acid molecules that encode these scFvs, and/or molecules.

[008] The present invention provides antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or polypeptide fragment of BLyS, said antibodies comprising, or alternatively consisting of, a polypeptide having the amino acid sequence of any one of the variable heavy ("VH") domains referred to in Table 1, below, or any one of the variable light ("VL") domains referred to in Table 1. In a preferred embodiment, antibodies of the present invention comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VH domain contained in SEQ ID NOS:1

-46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908, as referred to in Table 1 below. In another preferred embodiment, antibodies (including molecules comprising or alternatively consisting of, antibody fragments or variants thereof) of the present invention comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VL domain contained SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908, as referred to in Table 1 below. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies (e.g., including VH domains, VH CDRs, VL domains, or VL CDRs having an amino acid sequence of any one of those referred to in Table 1), that immunospecifically bind the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the soluble form and membrane-bound form of BLyS, are also encompassed by the invention, as are nucleic acid molecules that encode these antibodies, and/or molecules.

[009] The present invention also provides antibodies (including molecules comprising or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or a polypeptide fragment of BLvS, said antibodies comprising, or alternatively consisting of, a polypeptide having the amino acid sequence of any one of the VH domains referred to in Table 1, below, and any one of the VL domains referred to in Table 1. In a preferred embodiment, the antibodies of the invention comprise or alternatively consist of, a polypeptide having the amino acid sequence of a VH and VL domain contained in the same scFv referred to in Table 1. In another preferred embodiment, antibodies of the present invention, comprise, or alternatively consist of, a VH domain from an scFv of SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908, as disclosed in Table 1, and a VL domain from an scFv SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908, as disclosed in Table 1. In another preferred embodiment, antibodies of the present invention comprise, or alternatively consist of, the VH and VL domain from a single scFv of SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908, as disclosed in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies (e.g., including VH domains, VH CDRs, VL domains, or VL CDRs having an amino acid sequence of any one of those referred to in Table 1), that immunospecifically bind the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the soluble form and membrane-bound form of BLyS, are also encompassed by the invention,

as are nucleic acid molecules that encode these antibodies, and/or molecules.

[010] The present invention also provides antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or a polypeptide fragment of BLyS, said antibodies comprising, or alternatively consisting of, a polypeptide having the amino acid sequence of any one, two, three or more of the VH complementarity determining regions ("CDRs") (i.e., VH CDR1, VH CDR2, or VH CDR3) referred to in Table 1 and/or any one, two, three or more of the VL CDRs (i.e., VL CDR1, VL CDR2, or VL CDR3) referred to in Table 1. In one embodiment, antibodies of the present invention comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one of the VH CDR1s referred to in Table 1 and/or any one of the VL CDR1s referred to in Table 1. In another embodiment, antibodies of the present invention comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one of the VH CDR2s referred to in Table 1 and/or any one of the VL CDR2s referred to in Table 1. In a preferred embodiment, antibodies of the present invention comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one of the VH CDR3s referred to in Table 1 and/or any one of the VL CDR3s referred to in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies (e.g., including VH domains, VH CDRs, VL domains, or VL CDRs having an amino acid sequence of any one of those referred to in Table 1), that immunospecifically bind the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the soluble form and membrane-bound form of BLyS, are also encompassed by the invention, as are nucleic acid molecules that encode these antibodies, and/or molecules.

[011] In another embodiment, antibodies of the present invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) immunospecifically bind to a polypeptide or polypeptide fragment of BLyS, and comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one of the VH CDR1s referred to in Table 1, any one of the VH CDR2s referred to in Table 1. In another embodiment, antibodies of the present invention comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one of the VL CDR1s referred to in Table 1, any one of the VL CDR2s referred to in Table 1, any

in Table 1. In a preferred embodiment, antibodies of the present invention comprise, or alternatively consist of, at least one, two, three, four, five, six, or more CDRs that correspond to the same scFv referred to in Table 1, more preferably where CDR1, CDR2, and CDR3 of the VL domain correspond to the same scFv or where CDR1, CDR2, and CDR3 of the VH domain correspond to the same scFv, and most preferably where all six CDRs correspond to the same scFv referred to in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies (e.g., including VH domains, VH CDRs, VL domains, or VL CDRs having an amino acid sequence of any one of those referred to in Table 1), that immunospecifically bind the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the soluble form and membrane-bound form of BLyS, are also encompassed by the invention, as are nucleic acid molecules that encode these antibodies, and/or molecules.

The present invention also provides antibodies (including molecules [012]comprising, or alternatively consisting of, antibody fragments or variants thereof) that: immunospecifically bind to the soluble form of BLyS (e.g., a polypeptide consisting of amino acids 134 - 285 of SEO ID NO:3228); that immunospecifically bind to the membrane-bound form of BLyS (e.g., a polypeptide consisting of amino acids 1 - 285 of SEQ ID NO:3228 or a BLyS polypeptide expressed on the surface of monocytes) and/or that immunospecifically bind to both the soluble form and membrane-bound form of BLyS. In a preferred embodiment, antibodies of the present invention immunospecifically bind to the soluble form of BLyS and comprise, or alternatively consist of, a VH domain, VH CDR1, VH CDR2, VH CDR3, VL domain, VL CDR1, VL CDR2, and/or VL CDR3 corresponding to one or more scFvs, that immunospecifically bind to the soluble form of BLyS. In another preferred embodiment, antibodies of the present invention immunospecifically bind to the membrane-bound form of BLyS and comprise, or alternatively consist of, a VH domain, VH CDR1, VH CDR2, VH CDR3, VL domain, VL CDR1, VL CDR2, and/or VL CDR3 corresponding to one or more scFvs, that immunospecifically bind to the membrane-bound form of BLyS. In yet another preferred embodiment, antibodies of the present invention immunospecifically bind to the soluble form and membrane-bound form of BLyS and comprise, or alternatively consist of, a VH domain, VH CDR1, VH CDR2, VH CDR3, VL domain, VL CDR1, VL CDR2, and/or VL CDR3 corresponding to one or more scFvs, that immunospecifically binds to the soluble

form and membrane-bound form of BLyS. In another preferred embodiment, antibodies of the present invention comprise, or alternatively consist of, a VH domain and a VL domain corresponding to the same scFv disclosed in Table 1, which antibodies immunospecifically bind to the soluble form of BLyS, the membrane-bound form of BLyS, or both the soluble form and membrane-bound form of BLyS. Nucleic acid molecules encoding these antibodies are also encompassed by the invention. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies (e.g., including VH domains, VH CDRs, VL domains, or VL CDRs having an amino acid sequence of any one of those referred to in Table 1), that immunospecifically bind the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the soluble form and membrane-bound form of BLyS, are also encompassed by the invention, as are nucleic acid molecules that encode these antibodies, and/or molecules.

- [013] A VH domain of an amino acid sequence disclosed herein may be combined with
- [014] a VL domain of an amino acid sequence disclosed herein, or other VL domains, to provide a VH/VL pairing representing an antigen-binding site of an antibody. Similarly, a VL domain of an amino acid sequence disclosed herein may be combined with a VH domain of an amino acid sequence disclosed herein, or other VH domains. Further, one or more CDRs disclosed herein may be taken from a VH or VL domain and incorporated into a suitable framework as discussed *infra*.
- [015] The present invention provides antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof (including derivatives)) comprising, or alternatively consisting of, of VH domains, VL domains and/or CDRs described herein, which antibodies, immunospecifically bind to BLyS (e.g., soluble BLyS and membrane-bound BLyS) and can be routinely assayed for immunospecific binding to BLyS using methods known in the art, such as, for example, the immunoassays disclosed *infra*. Antibodies and antibody fragments or variants (including derivatives) of the invention may include, for example, one or more amino acid sequence alterations (addition, deletion, substitution and/or insertion of an amino acid residue). These alterations may be made in one or more framework regions and/or one or more CDR's. The antibodies of the invention (including antibody fragments, and variants and derivative thereof) can be routinely made by methods known in the art. Molecules

comprising, or alternatively consisting of, fragments or variants of any of the VH domains, VH CDRs, VL domains, and VL CDRs whose sequences are specifically disclosed herein may be employed in accordance with the present invention. Nucleic acid molecules encoding these antibodies and molecules (including fragments, variants, and derivatives) are also encompassed by the invention.

[016] The present invention also provides panels of antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants) wherein the panel members correspond to one, two, three, four, five, ten, fifteen, twenty, or more different antibodies of the invention (e.g., whole antibodies, Fabs, F(ab')2 fragments. Fd fragments, disulfide-linked Fvs (sdFvs), antiidiotypic (anti-Id) antibodies, and scFvs). The present invention further provides mixtures of antibodies, wherein the mixture corresponds to one, two, three, four, five, ten, fifteen, twenty, or more different antibodies of the invention (e.g., whole antibodies, Fabs, F(ab')2 fragments, Fd fragments, disulfide-linked Fvs (sdFvs), antiidiotypic (anti-Id) antibodies, and scFvs)). The present invention also provides for compositions comprising, or alternatively consisting of, one. two, three, four, five, ten, fifteen, twenty, or more antibodies of the present invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof). A composition of the invention may comprise, or alternatively consist of, one, two, three, four, five, ten, fifteen, twenty, or more amino acid sequences of one or more antibodies or fragments or variants thereof. Alternatively, a composition of the invention may comprise, or alternatively consist of, nucleic acid molecules encoding one or more antibodies of the invention.

[017] The present invention also provides for fusion proteins comprising an antibody (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) of the invention, and a heterologous polypeptide (i.e., a polypeptide unrelated to an antibody or antibody domain). Nucleic acid molecules encoding these fusion proteins are also encompassed by the invention. A composition of the present invention may comprise, or alternatively consist of, one, two, three, four, five, ten, fifteen, twenty or more fusion proteins of the invention. Alternatively, a composition of the invention may comprise, or alternatively consist of, nucleic acid molecules encoding one, two, three, four, five, ten, fifteen, twenty or more fusion proteins of the invention.

[018] The present invention also provides for a nucleic acid molecule, generally isolated, encoding an antibody (including molecules such as scFvs, which comprise, or alternatively consist of, an antibody fragment or variant thereof) of the invention. The present invention also provides a host cell transformed with a nucleic acid molecule of the invention and progeny thereof. The present invention also provides a method for the production of an antibody (including a molecule comprising, or alternatively consisting of, an antibody fragment or variant thereof) of the invention. The present invention further provides a method of expressing an antibody (including a molecule comprising, or alternatively consisting of, an antibody fragment or variant thereof) of the invention from a nucleic acid molecule. These and other aspects of the invention are described in further detail below.

[020] The present invention also encompasses methods and compositions for detecting, diagnosing and/or prognosing diseases or disorders associated with aberrant BLyS or BLyS receptor expression or inappropriate BLyS or BLyS receptor function in an animal, preferably a mammal, and most preferably a human, comprising using antibodies (including molecules which comprise, or alternatively consist of, antibody fragments or variants thereof) that immunospecifically bind to BLyS. Diseases and disorders which can be detected, diagnosed or prognosed with the antibodies of the invention include, but are not limited to, immune disorders (e.g., lupus, rheumatoid arthritis, multiple sclerosis, myasthenia gravis, Hashimoto's disease, and immunodeficiency syndrome), inflammatory disorders (e.g., asthma, allergic disorders, and rheumatoid arthritis), infectious diseases (e.g., AIDS), and proliferative disorders (e.g., leukemia, carcinoma, and lymphoma).

[021] In specific embodiments, the present invention encompasses methods and compositions for detecting, diagnosing and/or prognosing diseases or disorders associated with hypergammaglobulinemia (e.g., AIDS, autoimmune diseases, and some immunodeficiencies). In other specific embodiments, the present invention encompasses methods and compositions for detecting, diagnosing and/or prognosing diseases or disorders associated with hypogammaglobulinemia (e.g., an immunodeficiency).

[022] The present invention further encompasses methods and compositions for preventing, treating or ameliorating diseases or disorders associated with aberrant BLyS or BLyS receptor expression or inappropriate BLyS or BLyS receptor function in an animal, preferably a mammal, and most preferably a human, comprising administering to said

animal an effective amount of one or more antibodies (including molecules which comprise, or alternatively consist of, antibody fragments or variants thereof) that immunospecifically bind to BLyS. Diseases and disorders which can be prevented, treated or inhibited by administering an effective amount of one or more antibodies or molecules of the invention include, but are not limited to, immune disorders (e.g., lupus, rheumatoid arthritis, multiple sclerosis, myasthenia gravis, Hashimoto's disease, and immunodeficiency syndrome), inflammatory disorders (e.g., asthma, allergic disorders, and rheumatoid arthritis), infectious diseases (e.g., AIDS), and proliferative disorders (e.g., leukemia, carcinoma, and lymphoma).

[023] In specific embodiments, the present invention encompasses methods and compositions (e.g., antagonistic anti-BLyS antibodies) for preventing, treating or ameliorating diseases or disorders associated with hypergammaglobulinemia (e.g., AIDS, autoimmune diseases, and some immunodeficiency syndromes). In other specific embodiments, the present invention encompasses methods and compositions (e.g., agonistic anti-BLyS antibodies) for preventing, treating or ameliorating diseases or disorders associated with hypogammaglobulinemia (e.g., an immunodeficiency syndrome).

[024] Autoimmune disorders, diseases, or conditions that may be detected, diagnosed, prognosed, or monitored using the antibodies of the invention include, but are not limited to, autoimmune hemolytic anemia, autoimmune neonatal thrombocytopenia. idiopathic thrombocytopenia purpura, autoimmune neutropenia, autoimmunocytopenia, hemolytic anemia, antiphospholipid syndrome, dermatitis, gluten-sensitive enteropathy, allergic encephalomyelitis, myocarditis, relapsing polychondritis, rheumatic heart disease, glomerulonephritis (e.g., IgA nephropathy), Multiple Sclerosis, Neuritis, Uveitis Ophthalmia, Polyendocrinopathies, Purpura (e.g., Henloch-Scoenlein purpura), Reiter's Disease, Stiff-Man Syndrome, Autoimmune Pulmonary Inflammation, myocarditis, IgA glomerulonephritis, dense deposit disease, rheumatic heart disease, Guillain-Barre Syndrome, insulin dependent diabetes mellitis, and autoimmune inflammatory eye, autoimmune thyroiditis, hypothyroidism (i.e., Hashimoto's thyroiditis, systemic lupus erhythematosus, discoid lupus, Goodpasture's syndrome, Pemphigus, Receptor autoimmunities such as, for example, (a) Graves' Disease, (b) Myasthenia Gravis, and (c) insulin resistance, autoimmune hemolytic anemia, autoimmune thrombocytopenic purpura

, rheumatoid arthritis, schleroderma with anti-collagen antibodies, mixed connective tissue disease, polymyositis/dermatomyositis, pernicious anemia, idiopathic Addison's disease, infertility, glomerulonephritis such as primary glomerulonephritis and IgA nephropathy, bullous pemphigoid, Sjögren's syndrome, diabetes mellitus, and adrenergic drug resistance (including adrenergic drug resistance with asthma or cystic fibrosis), chronic active hepatitis, primary biliary cirrhosis, other endocrine gland failure, vitiligo, vasculitis, post-MI, cardiotomy syndrome, urticaria, atopic dermatitis, asthma, inflammatory myopathies, and other inflammatory, granulomatous, degenerative, and atrophic disorders).

[025] Immunodeficiencies that may be detected, diagnosed, prognosed, or monitored using the antibodies of the invention include, but are not limited to, severe combined immunodeficiency (SCID)-X linked, SCID-autosomal, adenosine deaminase deficiency (ADA deficiency), X-linked agammaglobulinemia (XLA), Bruton's disease, congenital agammaglobulinemia, X-linked infantile agammaglobulinemia, acquired agammaglobulinemia, adult onset agammaglobulinemia, late-onset agammaglobulinemia. dysgammaglobulinemia, hypogammaglobulinemia, transient hypogammaglobulinemia of infancy, unspecified hypogammaglobulinemia, agammaglobulinemia, common variable immunodeficiency (CVID) (acquired), Wiskott-Aldrich Syndrome (WAS), X-linked immunodeficiency with hyper IgM, non X-linked immunodeficiency with hyper IgM, selective IgA deficiency, IgG subclass deficiency (with or without IgA deficiency), antibody deficiency with normal or elevated Igs, immunodeficiency with thymoma, Ig heavy chain deletions, kappa chain deficiency, B cell lymphoproliferative disorder (BLPD), selective IgM immunodeficiency, recessive agammaglobulinemia (Swiss type), reticular dysgenesis, neonatal neutropenia, severe congenital leukopenia, thymic alymphoplasia-aplasia or dysplasia with immunodeficiency, ataxia-telangiectasia, short limbed dwarfism, X-linked lymphoproliferative syndrome (XLP), Nezelof syndromecombined immunodeficiency with Igs, purine nucleoside phosphorylase deficiency (PNP). MHC Class II deficiency (Bare Lymphocyte Syndrome) and severe combined immunodeficiency.

DEFINITIONS

[026] The term "antibody," as used herein, refers to immunoglobulin molecules and immunologically active portions of immunoglobulin molecules, i.e., molecules that

contain an antigen binding site that immunospecifically binds an antigen. As such, the term antibody encompasses not only whole antibody molecules, but also antibody fragments as well as variants (including derivatives) of antibodies and antibody fragments. Examples of molecules which are described by the term "antibody" in this application include, but are not limited to: single chain Fvs (scFvs), Fab fragments, Fab' fragments, F(ab')₂, disulfide linked Fvs (sdFvs), Fvs, and fragments comprising or alternatively consisting of, either a VL or a VH domain. The term "single chain Fv" or "scFv" as used herein refers to a polypeptide comprising a VL domain of antibody linked to a VH domain of an antibody. Antibodies that immunospecifically bind to BLyS may have cross-reactivity with other antigens. Preferably, antibodies that immunospecifically bind to BLyS can be identified, for example, by immunoassays or other techniques known to those of skill in the art, e.g., the immunoassays described in the Examples below.

[027] Antibodies of the invention include, but are not limited to, monoclonal, multispecific, human or chimeric antibodies, single chain antibodies, Fab fragments, F(ab') fragments, antiidiotypic (anti-Id) antibodies (including, e.g., anti-Id antibodies to antibodies of the invention), and epitope-binding fragments of any of the above. The immunoglobulin molecules of the invention can be of any type (e.g., IgG, IgE, IgM, IgD, IgA and IgY), class (e.g., IgG₁, IgG₂, IgG₃, IgG₄, IgA₁ and IgA₂) or subclass of immunoglobulin molecule.

[028] Preferably, an antibody of the invention comprises, or alternatively consists of, a VH domain, VH CDR, VL domain, or VL CDR having an amino acid sequence of any one of those referred to in Table 1, or a fragment or variant thereof.

[029] An antibody of the invention "which binds the soluble form of BLyS" is one which binds the 152 amino acid soluble form of the BLyS protein (amino acids 134-285 of SEQ ID NO:3228). In specific embodiments of the invention, an antibody of the invention "which binds the soluble form of BLyS" does not also bind the membrane-bound or membrane-associated form of BLyS. Assays which measure binding to the soluble form of BLyS include, but are not limited to, receptor binding inhibition assay or capture of soluble BLyS from solution as described in Examples 8 and 9.

[030] An antibody of the invention "which binds the membrane-bound form of BLyS" is one which binds the membrane-associated (uncleaved) BLyS protein. In

specific embodiments of the invention, an antibody of the invention "which binds the membrane-bound form of BLyS" does not also bind the soluble form of BLyS. Binding to HIS-tagged BLyS (as described herein) in an ELISA is an indicator that an antibody binds the membrane-bound form of BLyS, but should not be relied upon as proof of specificity for the membrane-bound form of BLyS. Assays that may be relied upon as proof of an antibody's specificity for membrane-bound BLyS, include, but are not limited to, binding to plasma membranes expressing BLyS as described in Example 2. An antibody of the invention "which binds the both the soluble form and the membrane-bound form of BLyS" is one which binds both the membrane-bound form and the soluble form of BLyS. [031] The term "variant" as used herein refers to a polypeptide that possesses a similar or identical function as a BLyS polypeptide, a fragment of BLyS, an anti-BLyS antibody or antibody fragment thereof, but does not necessarily comprise a similar or identical amino acid sequence of a BLyS polypeptide, a fragment of BLyS, an anti-BLyS antibody or antibody fragment thereof, or possess a similar or identical structure of a BLyS polypeptide, a fragment of BLyS, an anti-BLyS antibody or antibody fragment thereof. A variant having a similar amino acid refers to a polypeptide that satisfies at least one of the following: (a) a polypeptide comprising, or alternatively consisting of, an amino acid sequence that is at least 30%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95% or at least 99% identical to the amino acid sequence of a BLyS polypeptide, a fragment of BLyS, an anti-BLyS antibody or antibody fragment thereof (including a VH domain, VHCDR, VL domain, or VLCDR having an amino acid sequence of any one of those referred to in Table 1) described herein; (b) a polypeptide encoded by a nucleotide sequence, the complementary sequence of which hybridizes under stringent conditions to a nucleotide sequence encoding a BLyS polypeptide (e.g., SEQ ID NO:3228), a fragment of BLyS, an anti-BLyS antibody or antibody fragment thereof (including a VH domain, VHCDR, VL domain, or VLCDR having an amino acid sequence of any one of those referred to in Table 1), described herein, of at least 5 amino acid residues, at least 10 amino acid residues, at least 15 amino acid residues, at least 20 amino acid residues, at least 25 amino acid residues, at least 30 amino acid residues, at least 40 amino acid residues, at least 50 amino acid residues, at least 60 amino residues, at least 70 amino acid residues, at least 80 amino acid residues, at least 90 amino acid

residues, at least 100 amino acid residues, at least 125 amino acid residues, or at least 150 amino acid residues; and (c) a polypeptide encoded by a nucleotide sequence that is at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 99% or at least 99%, identical to the nucleotide sequence encoding a BLyS polypeptide, a fragment of BLyS, an anti-BLyS antibody or antibody fragment thereof (including a VH domain, VHCDR, VL domain, or VLCDR having an amino acid sequence of any one of those referred to in Table 1), described herein. A polypeptide with similar structure to a BLyS polypeptide, a fragment of BLyS, an anti-BLyS antibody or antibody fragment thereof, described herein refers to a polypeptide that has a similar secondary, tertiary or quarternary structure of a BLyS polypeptide, a fragment of BLyS, an anti-BLyS antibody, or antibody fragment thereof, described herein. The structure of a polypeptide can determined by methods known to those skilled in the art, including but not limited to, X-ray crystallography, nuclear magnetic resonance, and crystallographic electron microscopy.

To determine the percent identity of two amino acid sequences or of two nucleic acid sequences, the sequences are aligned for optimal comparison purposes (e.g., gaps can be introduced in the sequence of a first amino acid or nucleic acid sequence for optimal alignment with a second amino acid or nucleic acid sequence). The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in the first sequence is occupied by the same amino acid residue or nucleotide at the corresponding position in the second sequence, then the molecules are identical at that position. The percent identity between the two sequences is a function of the number of identical positions shared by the sequences (i.e., % identity = number of identical overlapping positions/total number of positions x 100%). In one embodiment, the two sequences are the same length.

[033] The determination of percent identity between two sequences can be accomplished using a mathematical algorithm known to those of skill in the art. An example of a mathematical algorithm for comparing two sequences is the algorithm of Karlin and Altschul *Proc. Natl. Acad. Sci. USA* 87:2264-2268(1990), modified as in Karlin and Altschul *Proc. Natl. Acad. Sci. USA* 90:5873-5877(1993). The BLASTn and BLASTx programs of Altschul, et al. *J. Mol. Biol.* 215:403-410(1990) have incorporated

such an alogrithm. BLAST nucleotide searches can be performed with the BLASTn program, score = 100, wordlength = 12 to obtain nucleotide sequences homologous to a nucleic acid molecules of the invention. BLAST protein searches can be performed with the BLASTx program, score = 50, wordlength = 3 to obtain amino acid sequences homologous to a protein molecules of the invention. To obtain gapped alignments for comparison purposes, Gapped BLAST can be utilized as described in Altschul et al. *Nucleic Acids Res.* 25:3389-3402(1997). Alternatively, PSI-BLAST can be used to perform an iterated search which detects distant relationships between molecules (*Id.*). When utilizing BLAST, Gapped BLAST, and PSI-BLAST programs, the default parameters of the respective programs (e.g., BLASTx and BLASTn) can be used. (*See* http://www.ncbi.nlm.nih.gov.)

[034] Another example of a mathematical algorithm utilized for the comparison of sequences is the algorithm of Myers and Miller, CABIOS (1989). The ALIGN program (version 2.0) which is part of the GCG sequence alignment software package has incorporated such an alogrithm. Other algorithms for sequence analysis known in the art include ADVANCE and ADAM as described in Torellis and Robotti Comput. Appl. Biosci., 10:3-5(1994); and FASTA described in Pearson and Lipman Proc. Natl. Acad. Sci. 85:2444-8(1988). Within FASTA, ktup is a control option that sets the sensitivity and speed of the search.

The term "derivative" as used herein, refers to a variant polypeptide of the invention that comprises, or alternatively consists of, an amino acid sequence of a BLyS polypeptide, a fragment of BLyS, or an antibody of the invention that immunospecifically binds to BLyS, which has been altered by the introduction of amino acid residue substitutions, deletions or additions. The term "derivative" as used herein also refers to a BLyS polypeptide, a fragment of BLyS, an antibody that immunospecifically binds to BLyS which has been modified, e.g., by the covalent attachment of any type of molecule to the polypeptide. For example, but not by way of limitation, a BLyS polypeptide, a fragment of BLyS, or an anti-BLyS antibody, may be modified, e.g., by glycosylation, acetylation, pegylation, phosphorylation, amidation, derivatization by known protecting/blocking groups, proteolytic cleavage, linkage to a cellular ligand or other protein, etc. A derivative of a BLyS polypeptide, a fragment of BLyS, or an anti-BLyS antibody, may be modified by chemical modifications using techniques known to those of

skill in the art, including, but not limited to, specific chemical cleavage, acetylation, formylation, metabolic synthesis of tunicamycin, etc. Further, a derivative of a BLyS polypeptide, a fragment of BLyS, or an anti-BLyS antibody, may contain one or more non-classical amino acids. A polypeptide derivative possesses a similar or identical function as a BLyS polypeptide, a fragment of BLyS, or an anti-BLyS antibody, described herein.

[036] The term "epitopes" as used herein refers to portions of BLyS having antigenic or immunogenic activity in an animal, preferably a mammal. An epitope having immunogenic activity is a portion of BLyS that elicits an antibody response in an animal. An eptiope having antigenic activity is a portion of BLyS to which an antibody immunospecifically binds as determined by any method known in the art, for example, by the immunoassays described herein. Antigenic epitopes need not necessarily be immunogenic.

amino acid sequence of at least 5 amino acid residues, at least 10 amino acid residues, at least 15 amino acid residues, at least 20 amino acid residues, at least 25 amino acid residues, at least 30 amino acid residues, at least 35 amino acid residues, at least 40 amino acid residues, at least 45 amino acid residues, at least 50 amino acid residues, at least 60 amino residues, at least 70 amino acid residues, at least 80 amino acid residues, at least 90 amino acid residues, at least 100 amino acid residues, at least 125 amino acid residues, at least 150 amino acid residues, at least 175 amino acid residues, at least 200 amino acid residues, or at least 250 amino acid residues, of the amino acid sequence of BLyS, or an anti-BLyS antibody (including molecules such as scFv's, that comprise, or alternatively consist of, antibody fragments or variants thereof) that immunospecifically binds to BLyS.

[038] The term "fusion protein" as used herein refers to a polypeptide that comprises, or alternatively consists of, an amino acid sequence of an anti-BLyS antibody of the invention and an amino acid sequence of a heterologous polypeptide (i.e., a polypeptide unrelated to an antibody or antibody domain).

[039] The term "host cell" as used herein refers to the particular subject cell transfected with a nucleic acid molecule and the progeny or potential progeny of such a cell. Progeny may not be identical to the parent cell transfected with the nucleic acid molecule due to mutations or environmental influences that may occur in succeeding

generations or integration of the nucleic acid molecule into the host cell genome.

DESCRIPTION OF THE FIGURES

- [040] Figure 1. ELISA results for three scFvs, I006E07, I008D05 and I016F04, that immunospecifically bind to U937 membranes, but not to bind to or cross-react with TNF-alpha or BSA.
- [041] Figure 2. The results for three scFvs, I016H07, I001C09 and I018D07, in a receptor inhibition assay.
- [042] Figure 3. ELISA results for two scFvs (I022D01 and I031F02) demonstrating their ability to bind to human BLyS and to cross-react with mouse BLyS, but not to bind to or cross-react with other antigens of the TNF ligand family.
- [043] Figure 4. ELISA results for three scFvs (I031F09, I050A12, and I051C04) binding to U937 plasma membranes when either BLyS or TNF-alpha is used as a competitor.
- [044] Figure 5. Kinetic analysis of scFv antibody I003C02. A dilution series of I003C02 from 3nM to 825nM is shown. Association and dissociation curves were generated using a BIAcore 2000 and BIAevaluation 3.0 software.
- [045] Figure 6. Typical titration curves for two scFv antibodies (I007F11 and I050A07) are shown in Figure 6. Unlabelled BLyS competed for binding to its receptor with an IC $_{50}$ value of 0.8 nM. The IC $_{50}$ values for I007F11 and I050A07 are 7.9 nM and 17.1 nM, respectively. The assay was performed in triplicate and standard error bars are shown.
- [001] Figure 7. ELISA results for three scFvs clones (I074B12, I075F12 and I075A02) that immunospecifically bind to immobilized BLyS, but not to U937 plasma membranes, TNF-alpha or BSA. As a control, a phage antibody that recognizes TNFα, is also shown in Figure 7.
- [047] Figure 8. The results for two scFvs (I025B09 and I026C04) in a receptor inhibition assay.
- [048] Figure 9. ELISA results for two scFvs clones (1067F05 and 1078D02) demonstrating their ability to bind to immobilized human BLyS and to cross-react with

immobilized mouse BLyS, but not to bind to or cross-react with other antigens of the TNF ligand family.

- [049] As a control, a phage antibody that recognizes TNF α , is also shown in Figure 7.
- [050] Figure 10. Kinetic analysis of scFV antibody I002A01. A dilution series of I002A01 from 3nM to 1650nM is shown. Association and dissociation curves were generated using a BIAcore 2000 and BIAevaluation 3.0 software.
- [051] Figure 11. Typical titration curves for two scFvs, I0068C06 and I074B12, are shown in Figure 11. Unlabelled BLyS competed for binding to its receptor with an inhibitory constant 50 (IC₅₀) value of 0.66 nM. The IC₅₀ values for I0068C06 and I074B12 are 61 nM and 13 nM, respectively. The assay was performed in triplicate and standard error bars are shown.
- [052] Figure 12. ELISA results for three clones (I079C01, I081C10 and I082A02) demonstrating their ability to bind histidine-tagged BLyS, U937 plasma membranes, but not to bind immobilized biotinylated BLyS.
- [053] Figure 13. ELISA results for three scFvs (I079B04, I079F08, and I080B01) binding to U937 plasma membranes when either histidine-tagged BLyS or biotinylated BLyS is used as a competitor.
- [054] Figure 14. An example of the dissociation section of a typical sensorgram for 8 scFvs is shown in Figure 14. An anti-TNF α antibody that does not recognize BLyS was included as a control. Of the 8 scFvs exemplified, I079F06 was identified for further study due to the relatively high numbers of RU's bound to the surface.
- [055] Figure 15. A typical example of the binding curves generated for the scFv antibody I082C03 is shown in Figure 15. The off-rate for this clone was calculated as $2x10^{-3}$ s⁻¹. The affinity of I082C03 was calculated as 20 nM, assuming 100% activity of the scFv.
- [056] Figure 16. ELISA results for three scFvs (I079B04, I079F08, and I080B01) binding to P388 plasma membranes when either histidine-tagged BLyS or biotinylated BLyS is used as a competitor.

DETAILED DESCRIPTION OF THE INVENTION

[057] The present invention encompasses antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS or a fragment or variant of BLyS. In particular, the invention provides antibodies such as, for example, single chain Fvs (scFvs) having an amino acid sequence of any one of SEQ ID NOS:1 - 2128, as referred to in Table 1. In particular, the present invention encompasses antibodies that immunospecifically bind to a polypeptide, a polypeptide fragment or variant, or an epitope of human BLyS (SEQ ID NOS:3228 and/or 3229) or BLyS expressed on human monocytes; murine BLyS (SEQ ID NOS:3230 and/or 3231) or BLyS expressed on murine monocytes; rat BLyS (either the soluble forms as given in SEQ ID NOS:3232, 3233, 3234 and/or 3235 or in a membrane associated form, e.g., on the surface of rat monocytes); or monkey BLyS (e.g., the monkey BLyS polypeptides of SEQ ID NOS:3236 and/or 3237, the soluble form of monkey BLyS, or BLyS expressed on monkey monocytes) (as determined by immunoassays known in the art for assaying specific antibody-antigen binding).

The polypeptide sequence shown in SEQ ID NO:3228 was obtained by sequencing and translating the cDNA of the HNEDU15 clone which was deposited on October 22, 1996 at the American Type Culture Collection, 10801 University Boulevard, Manassas, Virginia 20110-2209, and assigned ATCC Accession No. 97768. The deposited clone is contained in the pBluescript SK(-) plasmid (Stratagene, La Jolla, CA). The ATCC deposits were made pursuant to the terms of the Budapest Treaty on the international recognition of the deposit of microorganisms for the purposes of patent procedure.

[059] The polypeptide sequence shown in SEQ ID NO:3229 was obtained by sequencing and translating the cDNA of the HDPMC52 clone, which was deposited on December 10, 1998 at the American Type Culture Collection, and assigned ATCC Accession No. 203518. The deposited clone is contained in the pBluescript SK(-) plasmid (Stratagene, La Jolla, CA). The ATCC deposits were made pursuant to the terms of the Budapest Treaty on the international recognition of the deposit of microorganisms for the purposes of patent procedure.

[060] The BLyS polypeptides bound by the antibodies of the invention may be in monomers or multimers (i.e., dimers, trimers, tetramers and higher multimers).

Accordingly, the present invention relates to antibodies that bind monomers and multimers

of the BLyS polypeptides of the invention, their preparation, and compositions (preferably, pharmaceutical compositions) containing them. In specific embodiments, the antibodies of the invention bind BLyS monomers, dimers, trimers or tetramers. In additional embodiments, the antibodies of the invention bind at least dimers, at least trimers, or at least tetramers of BLyS.

homomers or heteromers. A BLyS homomer, refers to a multimer containing only BLyS polypeptides (including BLyS fragments, variants, and fusion proteins, as described herein). These homomers may contain BLyS polypeptides having identical or different amino acid sequences. In specific embodiments, the antibodies of the invention bind a BLyS homodimer (e.g., containing two BLyS polypeptides having identical or different amino acid sequences) or a BLyS homotrimer (e.g., containing three BLyS polypeptides having identical or different amino acid sequences). In a preferred embodiment, the antibodies of the invention bind homotrimers of BLyS. In additional embodiments, the antibodies of the invention bind a homomeric BLyS multimer which is at least a homodimer, at least a homotrimer, or at least a homotetramer.

[062]Heteromeric BLyS refers to a multimer containing heterologous polypeptides (i.e., polypeptides of a different protein) in addition to the BLyS polypeptides of the invention. In a specific embodiment, the antibodies of the invention bind a BLyS heterodimer, a heterotrimer, or a heterotetramer. In additional embodiments, the antibodies of the invention bind a heteromeric BLyS multimer which is at least a heterodimer, at least a heterotrimer, or at least a heterotetramer. In highly preferred embodiments, the antibodies of the invention bind a heterotrimer comprising both BLyS polypeptides and APRIL polypeptides (SEQ ID NO:3239; GenBank Accession No. AF046888; PCT International Publication Number WO97/33902; J. Exp. Med. 188(6):1185-1190) or fragments or variants thereof. In other highly preferred embodiments, the antibodies of the invention bind a heterotrimer comprising one BLyS polypeptide (including fragments or variants) and two APRIL polypeptides (including fragments or variants). In still other highly preferred embodiments, the antibodies of the invention bind a heterotrimer comprising two BLyS polypeptides (including fragments or variants) and one APRIL polypeptide (including fragments or variants). In a further

nonexclusive embodiment, the heteromers bound by the antibodies of the invention contain CD40 ligand polypeptide sequence(s), or biologically active fragment(s) or variant(s) thereof.

In particularly preferred embodiments, the antibodies of the invention bind homomeric, especially homotrimeric, BLyS polypeptides, wherein the individual protein components of the multimers consist of the mature form of BLyS (e.g., amino acids residues 134-285 of SEQ ID NO:3228, or amino acids residues 134-266 of SEQ ID NO:3229) or fragments or variants thereof. In other specific embodiments, antibodies of the invention bind heteromeric, especially heterotrimeric, BLyS polypeptides such as a heterotrimer containing two BLyS polypeptides and one APRIL polypeptide or a heterotrimer containing one BLyS polypeptide and two APRIL polypeptides, and wherein the individual protein components of the BLyS heteromer consist of the mature extracellular soluble portion of either BLyS (e.g., amino acids residues 134-285 of SEQ ID NO:3228, or amino acids residues 134-266 of SEQ ID NO:3229) or fragments or variants thereof, or the mature extracellular soluble portion APRIL (e.g., amino acid residues 105-250 of SEQ ID NO:3239) or fragments or variants thereof.

In specific embodiments, the antibodies of the invention bind conformational epitopes of a BLyS monomeric protein. In specific embodiments, the antibodies of the invention bind conformational epitopes of a BLyS multimeric, especially trimeric, protein. In other embodiments, antibodies of the invention bind conformational epitopes that arise from the juxtaposition of BLyS with a heterologous polypeptide, such as might be present when BLyS forms heterotrimers (e.g., with APRIL polypeptides (e.g., SEQ ID SEQ ID NO:3239)), or in fusion proteins between BLyS and a heterologous polypeptide.

[065] BLyS multimers bound by the antibodies of the invention may be the result of hydrophobic, hydrophilic, ionic and/or covalent associations and/or may be indirectly linked, by for example, liposome formation. Thus, in one embodiment, BLyS multimers, such as, for example, homodimers or homotrimers, are formed when polypeptides of the

invention contact one another in solution. In another embodiment, BLyS

heteromultimers, such as, for example, BLyS heterotrimers or BLyS heterotetramers, are formed when polypeptides of the invention contact antibodies to the polypeptides of the invention (including antibodies to the heterologous polypeptide sequence in a fusion protein of the invention) in solution. In other embodiments, BLyS multimers are formed by covalent associations with and/or between the BLyS polypeptides of the invention. Such covalent associations may involve one or more amino acid residues contained in the polypeptide sequence (e.g., that recited in SEQ ID NO:3228 or SEQ ID NO:3229). In one instance, the covalent associations are cross-linking between cysteine residues located within the polypeptide sequences which interact in the native (i.e., naturally occurring) polypeptide. In another instance, the covalent associations are the consequence of chemical or recombinant manipulation. Alternatively, such covalent associations may involve one or more amino acid residues contained in the heterologous polypeptide sequence in a BLyS fusion protein. In one example, covalent associations are between the heterologous sequence contained in a fusion protein (see, e.g., US Patent Number 5,478,925). In a specific example, the covalent associations are between the heterologous sequence contained in a BLyS-Fc fusion protein. In another specific example, covalent associations of fusion proteins of the invention are between heterologous polypeptide sequence from another TNF family ligand/receptor member that is capable of forming covalently associated multimers, such as for example, oseteoprotegerin (see, e.g., International Publication No. WO 98/49305, the contents of which are herein incorporated by reference in its entirety). In another specific example, covalent associations of fusion proteins of the invention are between heterologous polypeptide sequence from CD40L, or a soluble fragment thereof. In another embodiment, two or BLyS polypeptides are joined through synthetic linkers (e.g., peptide, carbohydrate or soluble polymer linkers). Examples include those peptide linkers described in U.S. Pat. No. 5,073,627 (hereby incorporated by reference). Proteins comprising multiple BLyS polypeptides separated by peptide linkers may be produced using conventional recombinant DNA technology. [066] In one embodiment, antibodies of the invention immunospecifically bind a BLyS polypeptide having the amino acid sequence of SEQ ID NO:3228 or as encoded by

the cDNA clone contained in ATCC No. 97768, or a polypeptide comprising a portion

(i.e., a fragment) of the above polypeptides. In another embodiment, the invention

provides an antibody that binds an isolated BLyS polypeptide having the amino acid sequence of SEQ ID NO:3229 or the amino acid sequence encoded by the cDNA clone contained in ATCC No. 203518, or a an antibody that binds polypeptide comprising a portion (i.e, fragment) of the above polypeptides.

[067] Antibodies of the present invention immunospecifically bind to polypeptides comprising or alternatively, consisting of, the amino acid sequence of SEQ ID NO:3228, encoded by the cDNA contained in the plasmid having ATCC accession number 97768, or encoded by nucleic acids which hybridize (e.g., under stringent hybridization conditions) to the nucleotide sequence contained in the deposited clone. Antibodies of the present invention also bind to fragments of the amino acid sequence of SEQ ID NO:3228, encoded by the cDNA contained in the plasmid having ATCC accession number 97768, or encoded by nucleic acids which hybridize (e.g., under stringent hybridization conditions) to the nucleotide sequence contained in the deposited clone.

[068] Additionally, antibodies of the present invention bind polypeptides comprising or alternatively, consisting of, the amino acid sequence of SEQ ID NO:3229, encoded by the cDNA contained in the plasmid having ATCC accession number 203518, or encoded by nucleic acids which hybridize (e.g., under stringent hybridization conditions) to the nucleotide sequence contained in the deposited clone. Antibodies of the present invention also bind to fragments of the amino acid sequence of SEQ ID NO:3229, encoded by the cDNA contained in the plasmid having ATCC accession number 203518, or encoded by nucleic acids which hybridize (e.g., under stringent hybridization conditions) to the nucleotide sequence contained in the deposited clone.

[069] In addition, antibodies of the invention bind polypeptides or polypeptide fragments comprising or alternatively, consisting of, an amino acid sequence contained in SEQ ID NOS: 3230 through 3237.

[070] In specific embodiments, the antibodies of the present invention immunospecifically bind polypeptide fragments including polypeptides comprising or alternatively, consisting of, an amino acid sequence contained in SEQ ID NO:3228, encoded by the cDNA contained in the deposited clone, or encoded by nucleic acids which hybridize (e.g., under stringent hybridization conditions) to the nucleotide sequence contained in the deposited clone. Protein fragments may be "free-standing," or comprised

within a larger polypeptide of which the fragment forms a part or region, most preferably as a single continuous region. Representative examples of polypeptide fragments that may be bound by the antibodies of the present invention, include, for example, fragments that comprise or alternatively, consist of from about amino acid residues: 1 to 50, 51 to 100, 101 to 150, 151 to 200, 201 to 250, and/or 251 to 285 of SEQ ID NO:3228. Moreover, polypeptide fragments can be at least 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 175 or 200 amino acids in length.

- [071] In specific embodiments, antibodies of the present invention bind polypeptide fragments comprising, or alternatively consisting of, amino acid residues: 1-46, 31-44, 47-72, 73-285, 73-83, 94-102, 148-152, 166-181, 185-209, 210-221, 226-237, 244-249, 253-265, and/or 277-285 of SEQ ID NO:3228.
- [072] It will be recognized by one of ordinary skill in the art that mutations targeted to regions of a BLyS polypeptide of SEQ ID NO:3228 which encompass the nineteen amino acid residue insertion which is not found in the BLyS polypeptide sequence of SEQ ID NO:3229 (i.e., amino acid residues Val-142 through Lys-160 of the sequence of SEQ ID NO:3229) may affect the observed biological activities of the BLyS polypeptide. More specifically, a partial, non-limiting and non-exclusive list of such residues of the BLyS polypeptide sequence which may be targeted for mutation includes the following amino acid residues of the BLyS polypeptide sequence as shown in SEQ ID NO:3228: V-142; T-143; Q-144; D-145; C-146; L-147; Q-148; L-149; I-150; A-151; D-152; S-153; E-154; T-155; P-156; T-157; I-158; Q-159; and K-160. Thus, in specific embodiments, antibodies of the present invention that bind BLyS polypeptides which have one or more mutations in the region from V-142 through K-160 of SEQ ID NO:3228 are contemplated.
- [073] Polypeptide fragments may be "free-standing," or comprised within a larger polypeptide of which the fragment forms a part or region, most preferably as a single continuous region. Representative examples of polypeptide fragments that may be bound by antibodies of the present invention, include, for example, fragments that comprise or alternatively, consist of from about amino acid residues: 1 to 15, 16-30, 31-46, 47-55, 56-72, 73-104, 105-163, 163-188, 186-210 and 210-284 of the amino acid sequence disclosed in SEQ ID NO:3228. Additional representative examples of polypeptide fragments that may be bound by antibodies of the present invention, include, for example, fragments that

comprise or alternatively, consist of from about amino acid residues: 1 to 143, 1-150, 47-143, 47-150, 73-143, 73-150, 100-150, 140-145, 142-148, 140-150, 140-200, 140-225, and 140-266 of the amino acid sequence disclosed in SEQ ID NO:3229. Moreover, polypeptide fragments that may be bound by antibodies of the present invention, can be at least 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 175 or 200 amino acids in length. In this context, "about" means the particularly recited ranges and ranges larger or smaller by several, a few, 5, 4, 3, 2 or 1 amino acid residues at either or both the amino- and carboxy-termini.

[074] Additional preferred embodiments encompass antibodies that bind polypeptide fragments comprising, or alternatively consisting of, the predicted intracellular domain of BLyS (e.g., amino acid residues 1-46 of SEQ ID NO:3228), the predicted transmembrane domain of BLyS (e.g., amino acid residues 47-72 of SEQ ID NO:3228), the predicted extracellular domain of BLyS (e.g., amino acid residues 73-285 of SEQ ID NO:3228), the mature soluble extracellular domain of BLyS (e.g., amino acids residues 134-285 of SEQ ID NO:3228), the predicted TNF conserved domain of BLyS (e.g., amino acids 191 to 284 of SEQ ID NO:3228), and a polypeptide comprising, or alternatively, consisting of the predicted intracellular domain fused to the predicted extracellular domain of BLyS (amino acid residues 1-46 fused to amino acid residues 73-285 of SEQ ID NO:3228).

[075] Further additional preferred embodiments encompass polypeptide fragments comprising, or alternatively consisting of, the predicted intracellular domain of BLyS (amino acid residues 1-46 of SEQ ID NO:3229), the predicted transmembrane domain of BLyS (amino acid residues 47-72 of SEQ ID NO:3229), the predicted extracellular domain of BLyS (amino acid residues 73-266 of SEQ ID NO:3229), the predicted TNF conserved domain of BLyS (amino acids 172 to 265 of SEQ ID NO:3229), and a polypeptide comprising, or alternatively, consisting of the predicted intracellular domain fused to the predicted extracellular domain of BLyS (amino acid residues 1-46 fused to amino acid residues 73-266 of SEQ ID NO:3229).

[076] Certain additional embodiments of the invention encompass antibodies that bind polypeptide fragments comprising, or alternatively consisting of, the predicted betapleated sheet regions of the BLyS polypeptides of SEQ ID NO:3228 and SEQ ID NO:3229. These polypeptide fragments comprising the beta-pleated sheets of BLyS

comprise, or alternatively consist of, amino acid residues Gln-144 to Ala-151, Phe-172 to Lys-173, Ala-177 to Glu-179, Asn-183 to Ile-185, Gly-191 to Lys-204, His-210 to Val-219, Leu-226 to Pro-237, Asn-242 to Ala-251, Gly-256 to Ile-263 and/or Val-276 to Leu-284 of SEQ ID NO:3228. In another, nonexclusive embodiment, these polypeptide fragments comprising the beta-pleated sheets of BLyS comprise, or alternatively consist of, amino acid residues Phe-153 to Lys-154, Ala-158 to Glu-160, Asn-164 to Ile-166, Gly-172 to Lys-185, His-191 to Val-200, Leu-207 to Pro-218, Asn-223 to Ala-232, Gly-237 to Ile-244 and/or Val-257 to Leu-265 of SEQ ID NO:3229.

A partial, non-limiting, and exemplary list of polypeptides that may be [077]bound by the antibodies of the invention includes polypeptides that comprise, or alternatively consist of, combinations of amino acid sequences of the invention includes, for example, [Met-1 to Lys-113] fused to [Leu-114 to Thr-141] fused to [Val-142 to Lys-160] fused to [Gly-161 to Gln-198] fused to [Val-199 to Ala-248] fused to [Gly-249 to Leu-285] of SEQ ID NO:3228; or [Met-1 to Lys-113] fused to [Val-142 to Lys-160] fused to [Gly-161 to Gln-198] fused to [Val-199 to Ala-248] fused to [Gly-249 to Leu-285] of SEQ ID NO:3228; or [Met-1 to Lys-113] fused to [Leu-114 to Thr-141] fused to [Val-142] to Lys-160] fused to [Gly-161 to Gln-198] fused to [Gly-249 to Leu-285] of SEQ ID NO:3228. Other combinations of amino acids sequences that may be bound by the antibodies of the invention may include the polypeptide fragments in an order other than that recited above (e.g., [Leu-114 to Thr-141] fused to [Val-199 to Ala-248] fused to [Gly-249 to Leu-285] fused to [Val-142 to Lys-160] of (SEQ ID NO:3228). Other combinations of amino acids sequences that may be bound by the antibodies of the invention may also include heterologous polypeptide fragments as described herein and/or other polypeptides or polypeptide fragments of the present invention (e.g., [Met-1 to Lys-113] fused to [Leu-114 to Thr-141] fused to [Val-142 to Lys-160] fused to [Gly-161 to Gln-198] fused to [Gly-249 to Leu-285] of SEQ ID NO:3228 fused to a FLAG tag; or [Met-1 to Lys-113] of SEQ ID NO:3228 fused to [Leu-114 to Thr-141] of SEQ ID NO:3228 fused to [Glu-135 to Asn-165] of SEQ ID NO:39 fused to [Val-142 to Lys-160] of SEQ ID NO:3228 fused to [Gly-161 to Gln-198] of SEQ ID NO:3228 fused to [Val-199] to Ala-248] of SEQ ID NO:3228 fused to [Gly-249 to Leu-285] of SEQ ID NO:3228).

[078] A partial, non-limiting, and exemplary list of polypeptides that may be bound by the antibodies of the invention includes polypeptides that comprise, or

alternatively consist of, combinations of amino acid sequences includes, for example, [Met-1 to Lys-113] fused to [Leu-114 to Thr-141] fused to [Gly-142 to Gln-179] fused to [Val-180 to Ala-229] fused to [Gly-230 to Leu-266] of SEQ ID NO:3229; [Met-1 to Lys-113] fused to [Gly-142 to Gln-179] fused to [Val-180 to Ala-229] fused to [Gly-230 to Leu-266] of SEQ ID NO:3229; or [Met-1 to Lys-113] fused to [Leu-114 to Thr-141] fused to [Gly-142 to Gln-179] fused to [Gly-230 to Leu-266] of SEQ ID NO:3229. Other of amino acids sequences that may be bound by the antibodies of the invention combinations may include the polypeptide fragments in an order other than that recited above (e.g., [Leu-114 to Thr-141] fused to [Val-180 to Ala-229] fused to [Gly-230 to Leu-266] fused to [Gly-142 to Gln-179] of SEQ ID NO:3229). Other combinations of amino acid sequences that may be bound by the antibodies of the invention may also include heterologous polypeptide fragments as described herein and/or other polypeptides or polypeptide fragments of the present invention (e.g., [Met-1 to Lys-113] fused to [Leu-114 to Thr-141] fused to [Gly-142 to Gln-179] fused to [Gly-230 to Leu-266] of SEO ID NO:3229 fused to a FLAG tag (SEQ ID NO:3238) or, [Met-1 to Lys-113] of SEQ ID NO:3229 fused to [Leu-114 to Thr-141] of SEQ ID NO:3229 fused to [Glu-135 to Asn-165] of SEQ ID NO:39 fused to [Gly-142 to Gln-179] of SEQ ID NO:3229 fused to [Val-180 to Ala-229] of SEQ ID NO:3229 fused to [Gly-230 to Leu-266] of SEO ID NO:3229. [079] Additional embodiments of the invention encompass antibodies that bind BLyS polypeptide fragments comprising, or alternatively consisting of, functional regions of polypeptides of the invention, such as the Garnier-Robson alpha-regions, beta-regions, turn-regions, and coil-regions, Chou-Fasman alpha-regions, beta-regions, and coil-regions, Kyte-Doolittle hydrophilic regions and hydrophobic regions, Eisenberg alpha- and beta-amphipathic regions, Karplus-Schulz flexible regions, Emini surface-forming regions and Jameson-Wolf regions of high antigenic index set out in Tables 9 and 10 and as described herein. In a preferred embodiment, the polypeptide fragments bound by the antibodies of the invention are antigenic (i.e., containing four or more contiguous amino acids having an antigenic index of greater than or equal to 1.5, as identified using the

[080] The data representing the structural or functional attributes of the BLyS polypeptide of SEQ ID NO:3228 (Table 9) or the BLyS polypeptide of SEQ ID NO:3229

default parameters of the Jameson-Wolf program) of a complete (i.e., full-length) BLyS

polypeptide (e.g., SEQ ID NOS:3228 and 3229).

(Table 10), as described above, was generated using the various modules and algorithms of the DNA*STAR set on default parameters. Column I represents the results of a Garnier-Robson analysis of alpha helical regions; Column III represents the results of a Chou-Fasman analysis of alpha helical regions; Column IV represents the results of a Garnier Robson analysis of beta sheet regions; Column IV represents the results of a Chou-Fasman analysis of turn regions; Column VI represents the results of a Garnier Robson analysis of turn regions; Column VII represents the results of a Garnier Robson analysis of turn regions; Column VII represents the results of a Garnier Robson analysis of coil regions; Column VIII represents a Kyte-Doolittle hydrophilicity plot; Column IX represents a Hopp-Woods hydrophobicity plot; Column X represents the results of an Eisenberg analysis of alpha amphipathic regions; Column XII represents the results of a Karplus-Schultz analysis of flexible regions; Column XIII represents the Jameson-Wolf antigenic index score; and Column XIV represents the Emini surface probability plot.

[081] In a preferred embodiment, the data presented in columns VIII, IX, XIII, and XIV of Tables 9 and 10 can be used to determine regions of the BLyS polypeptide of SEQ ID NO:3228 (Table 9) or the BLyS polypeptide of SEQ ID NO:3229 (Table 10) which exhibit a high degree of potential for antigenicity. Regions of high antigenicity are determined from the data presented in columns VIII, IX, XIII, and/or XIV by choosing values which represent regions of the polypeptide which are likely to be exposed on the surface of the polypeptide in an environment in which antigen recognition may occur in the process of initiation of an immune response.

[082] The above-mentioned preferred regions set out in Tables 9 and 10 include, but are not limited to, regions of the aforementioned types identified by analysis of the amino acid sequence set out in SEQ ID NO:2. As set out in Tables 9 and 10, such preferred regions include Garnier-Robson alpha-regions, beta-regions, turn-regions, and coil-regions, Chou-Fasman alpha-regions, beta-regions, and turn-regions, Kyte-Doolittle hydrophilic regions, Eisenberg alpha- and beta-amphipathic regions, Karplus-Schulz flexible regions, Jameson-Wolf regions of high antigenic index and Emini surface-forming regions. Preferably, antibodies of the present invention bind BLyS polypeptides or BLyS polypeptide fragments and variants comprising regions of BLyS that combine several

structural features, such as several (e.g., 1, 2, 3, or 4) of the same or different region features set out above and in Tables 9 and 10.

Res Po	osition	1.	п	ш	IV .	\mathbf{v}	VI	VII	VIII	IX	X	XI	ж	XIII	XIV
Met	1	A							0.73	-0.71				0.95	1.39
Asp	2	Α					T		1.12	-0.66				1.15	1.56
Asp	3	A					T		1.62	-1.09				1.15	2.12
Ser	4	Ā			-		T		2.01	-1.51		į	• •	1.15	4.19
Thr	5	A	·				T		2.40	-2.13			F	1.30	4.35
Glu	6	Ā	À	Ţ					2.70	-1.73	•		F	0.90	4.51
Arg	7	A	Ā		·				2.81	-1.34	*		F	0.90	4.51
Glu	8	A	Ā	·	·	·			2.00	-1.73	*		F	0.90	6.12
Gin	9.	A	A	•		·	·		1.99	-1.53	*		F	0.90	2.91
Ser	10	Ä	••	•	В	•	•	•	2.00	-1.04	*		F	0.90	2.15
Arg	11	Ä	•	•,	B	•	•	•	1.33	-0.66	*		F	0.90	1.66
Leu	12	· A	:	•	·B	•	• .	•	0.41	-0.09	*		F	0.45	0.51
Thr	13	Ā	•	•	В	•	•	•	0.46	0.20			F	-0.15	0.31
Ser	14	Â	A	•				•	0.50	-0.19				0.30	0.32
Cys	15	Â	Â.	. •	•	•	•	•	0.91	-0.19			•	0.30	0.32
Leu	16	A	A	•	•	•	•		0.80	-0.19	*	*	F	0.90	1.06
Lys	17	Â	Ā	•	•	•	•	•	1.61	-1.36			F	0.90	1.37
-	18	Ā	Ā	. •	•	•	•	•	1.32	-1.74	. '	•	F	0.90	4.44
Lys	19	Â	Â	•	•	•	•	•	1.67	-1.70	•		F	0.90	5.33
Arg Glu	20	A	A	•	•	•	•	•	1.52	-2.39	•		F	0.90	
				•	•	•	•	•	2.38	-2.39	•		F		5.33
Glu	. 21	A	A	•	•	•	•	•	2.33	-1.70	•		F	0.90	2.20
Met	22	A A	A A	•	•	•	• ,	• 4	1.62				F	0.90	2.24
Lys	23			•	•	•	•	•	0.66	-1.70			F	0.90	2.24
Leu	24	A	A	. •	•	•	•	•		-1.13	. *			0.75	0.69
Lys	25	A	A	•		•	•	•	0.36	-0.49	:	•	F	0.45	0.52
Glu	26	A	A	•	В	•	•	•	-0.53	-0.71		I	. •	0.60	0.35
Cys	27	A	- A	•	В	•	•	•	-0.74	-0.03	*	-	•	0.30	0.30
Val	28	A	A	•	В	•	•	•	-1.00	-0.03			•	0.30	0.12
Ser	29	A	A	•	В	•	•	•	-0.08	0.40	*	-	•	-0.30	0.11
Ile	30	A	•	• •	В	•	•	•	-0.08	0.40	•	-	•	-0.30	0.40
Leu	- 31	A	•	•	В	•	•		-0.08	-0.17		•	:	0.45	1.08
Pro	32	•	•	•	В		. •	С	0.29	-0.81	•	•	F	1.10	1.39
Arg	33	•	٠.	•	•	T	•	•	0.93	-0.81	•	•	F	1.50	2.66
Lys	34	•	•	•	• •	T			0.93	-1.07		:	F	1.84	4.98
Glu	35	•	•	•	•	•		C	0.97	-1.37		-	F	1.98	4.32
Ser	36	•	•	• .	•	•	T	C	1.89	-1.16		•	F	2.52	1.64
Pro	37	•	•	•	• .	<u>.</u>	T	С	1.80	-1.16		•	F	2.86	1.60
Ser	38	•	•	•	•	T	T	•	1.39	-0.77	•	• .	F	3.40	1.24
Val	39	A	٠.	•	•	•	T	•	1.39	-0.39	:	7	F	2.36	1.24
Arg	40	Ą	• .	. •	• '	•	•	•	1.39	-0.77	•	•	F	2.46	1.60
Ser	41	A	•	•	•	•	<u>.</u>	•	1.34	-1.20	•	•	F	2.46	2.00
Ser	42	•	•	•	•	T	T	•	1.60	-1.16	•		F	3.06	2.67
Lys	43	•	•	•	•	T	T	•	1.09	-1.80	•	*	F	3.06	2.72
Asp	44	• .	•	•	•	T	T .	•	1.13	-1.11			F	3.40	1.67
Gly	45	Α.,	•	•	• .	•	T	• *	0.43	-0.81	*	*	F	2.66	1.03
Lys	46	A	A	•	•	•	•	•	0:14	-0.70	•	•	F.	1.77	0.52
Leu	47	A	A	• .	•	•	•	•	0.13	-0.20	*	•	•	0.98	0.31
Leu	48	A	A	•		•		•	-0.72	0.29	. •	•		0.04	0.46
Ala	49	A	A	•			•	•	-1.53	0.54		*	•	-0.60	0.19
Ala	50	Α	A	•				•	-2.00	1.23		•		-0.60	0.19

Table 9 (continued)

Res P	osition	1	. , u	m	IV	v	VI	VII	VIII.	IX	x	x	XII	ХШ	XIV
Thr	51	A	Α	•					-2.63	1.23	•			-0.60	0.19
Leu	52	A	A	•	•		•	•	-2.63	1.04	•			-0.60	0.19
Leu	- 53	A	A		•	•	•	•	-2.63	1.23	•			-0.60	0.15
Leu	54	Α	A	•	•		•		-2.34	1.41				-0.60	0.09
Ala	55	Α	A	•	٠.		•		-2.42	1.31				-0.60	0.14
Leu	56	Α	Α			•			-2.78	1.20		•		-0.60	0.09
Leu	57	Α		, .	•		Τ.		-2.78	1.09	` . .			-0.20	0.06
Ser	58	A			•	•	T		-2.28	1.09				-0.20	0.05
Cys	59	A					T		-2.32	1.07				-0.20	0.09
Cys	60	A				•	T	•	-2.59	1.03				-0.20	0.08
Leu .	61	•		·B	В .			. •	-2.08	0.99	:			-0.60	0.04
Thr	62			В	В			•	-1.97	0.99				-0.60	0.11
Val	63		. •	В	В				-1.91	1.20			•	-0.60	0.17
Val	64		•	В.	В	• .			-1.24	1.39				-0.60	0.33
Ser	65			В	В	•			-1.43	1.10		•		-0.60	0.40
Phe	. 66	Α			В		• •	•	-1.21	1.26				-0.60	0.40
Tyr	67	. A	. •		В				-1.49	1.11				-0.60	0.54
Gln	68	Α	٠.		В	•			-1.44	0.97				-0.60	0.41
Val	69	A	· .		В.				-0.59	1.27				-0.60	0.39
Ala	70	· A			В				-0.63	0.89				-0.60	0.43
Ala	- 71	A			В				0.07	0.56				-0.60	0.25
Leu	72	Α		:			T		-0.50	0.16		. •		0.10	0.55
Gln	73	A				•	T		-1.09	0.20			F	0.25	0.45
Gly	74	Α			•		T		-0.53	0.20			F	0.25	0.45
Asp	75 ·	A					T		-0.76	0.09			F	0.25	0.73
Leu	76	A	A.				٠.		-0.06	0.09		•	F	-0.15	0.35
Ala	77	· A	Α						0.17	-0.31	į	. *	•	0.30	0.69
Ser	78	A	A				: .		0.17	-0.24			·	0.30	0.42
Leu	79	A 5	A			٠			-0.30	-0.24		*	•	0.30	0.88
Arg	80	A	Α						-0.30	-0.24	·	*		0.30	0.72
Ala	81	A	Α						0.17	-0.34	Ĭ		•	0.30	0.93
Glu	82	. A	A						0.72	-0.30	·	*	•	0.45	1.11
Leu	83	A	A			•			0.99	-0.49	•	*	•	0.30	0.77
Gln	84	A	Α	•					1.21	0.01	•		•	-0.15	1.04
Gly	85	Α	Α						1.10	0.01			•	-0.30	0.61
His	86	A	Α				_		1.73	0.01	•	*	•	-0.15	1.27
His	87·	A	A						0.92	-0.67		*	·	0.75	1.47
Ala	. 88	A	A						1.52	-0.39				0.45	1.22
Glu	. 89	Α	A						0.93	-0.39			•	0.45	1.39
Lys	90	. A	A						0.93	-0.39			F	0.60	1.03
Leu	91	. A					T		0.38	-0.46		·	•	0.85	1.01
Pro ·	92	Α					T		0.07	-0.46				0.70	0.59
Ala	93	A					T		0.07	-0.03			- [-	0.70	0.29
Gly	94	Α	•				T		-0.14	0.47	·		·	-0.20	0.36
Ala	95	A		•					0.14	0.21			•	-0.10	0.36
Gly	96	A				-	-	-	0.08	-0.21	•		F	0.65	0.30
Ala	97	A				Ē		•	-0.06	-0.21	•	•	F	0.65	0.71
Pro	98	Ā	•			•	•	•	-0.28	-0.21	•	•	F	0.65	0.72
Lys	99	Ā	A			•	•	•	0.07	-0.03	•		F	0.45	
Ala	100	A	A	•		• .	•	•	0.66	-0.03 -0.46	•	• •	F	0.60	0.59
	-30	43	. 15	•	• .	•	•	•	V.00	~0.40	•.	•	r	U.OU	1.01

Table 9 (continued)

Res P	osition	I.	п	ш	IV .	v .	VI	· VII	VIII	IX	x	IX	XII	XIII	XIV
Gly	101	A	A			•			0.41	-0.96			F	0.90	1.13
Leu	102	Α	Α.						0.79	-0.89			F	0.75	0.57
Glu	103	A	A	• •					0.41	-0.46			F	0.45	0.88
Glu	104	Α	· A						-0.49	-0.46			·F	0.45	0.89
Ala	105	Α	A			•.			-0.21	-0.24		•	•	0.30	0.81
Pro	106	Α	A	•					-0.46	-0.44	·		•	0.30	0.67
Ala	107	Α	Α					_	0.01	0.06	·	•	•	-0.30	0.39
Val	108	A	Α						-0.80	0.49	•	•	•	-0.60	0.38
Thr	109	· A	A						-0.76	0.67	•	* ,	• .	-0.60	0.20
Ala	110	Α	A						-1.06	0.24	•		•	-0.30	0.40
Gly	111	A	Α						-1.54	0.43			•	-0.60	0.38
Leu	112	. A	A						-0.96	0.57	*		•	-0.60	0.38
Lys	113		A	В	_		· ·	·	-0.31	0.09			•	-0.30	0.25
Ile	114		A	В	:				-0.21	0.01	*		•	-0.30	0.59
Phe	115		A	. B		•	•	• •	-0.21	0.01		•	•	0.15	
Glu	116	•	Ā		•	•	•	c	-0.08	-0.17		•	F	1.25	1.15
Pro	117	•	Ä	•	•	•	•	č	0.39	0.26		:	F	1.10	0.58 1.28
Pro	118	•	••	•	• .	• .	•	č	0.34	-0.00			F		
Ala	119	• .	•	. •	•	•	Ť	Ċ.	0.89	-0.79	•			2.20	1.47
Pro	120	. •	•	•	•	•	Ť	C	1.59	-0.79	•		F	3.00	1.47
Gly	121	•	• ·	•	•	T	Ť	_	1.29		•	•	F	2.25	0.94
Glu	122	. •	•	•	•	T	Ť	•		-0.39	••	•	F.	2.15	0.98
Gly	123	•	•	•	•			C	1.20 1.41	-0.43	•	•	F	2.00	1.30
Asn	124	•		•	•	•	· †	-	-	-0.54	•	•	F	1.60	1.12
Ser	125	•	•	. •	•	•	T	C C	2.00	-0.57	. •	:	E.	1.50	1.97
Ser	126	. •	•	•	• .	•	T	C	1.91	-0.60	•	- :	F	1.50	1.82
Gln	120		•	•	•	•			2.37	-0.21	•	-	F	1.54	2.47
Asn	127	•	•	•	•	•	T	C	2.37	-0.64	•	•	F	2.18	3.01
	128	•	•	•	•	• .	<u>.</u> .	C	2.76	-0.64	•	•	F	2.32	3.61
Ser		•	•	•	• .		T	С	2.87	-1.03	:	• •	F	2.86	5.39
Arg	130	•	٠.	• ·	•	T	Ţ	•	2.58	-1.41	•	, •	· F	3.40	6.09
Asn	131	•	•	•	•	T	T	•	2.02	-1.31		•	F	3.06	3.83
Lys			•	•	•	T	T	•	2.02	-1.07	•	•	F	2.72	2.12
Arg	133	•	•	•	• •	T	•		1.68	-1.06	*		F	2.18	1.88
Ala	134	•	•	•	• '	• .	•	C	1.77	-0.63	•	•	F	1.64	1.15
Val	135	•	•	•	•	.•	•	C	1.66	-0.60	•	•	F	1.49	0.89
Gln	136		•	• .	•	. •	÷	C	1.66	-0.60			F	1.83	0.79
Gly	137	. •	•	•	• .	•	T	C	1.30	-0.60	*	•	F	2.52	1.35
Pro	138	•	•	•	•	<u>.</u>	T	С	0.33	-0.61		•	F	2.86	2.63
Glu	139	•	• • .	•	• .	T	T	:	0.61	-0.61			. F	3.40	1.13
Glu	140	A.	• .	•	•	•	T	•	1.47	-0.53	•	•	F	2.66	1.64
Thr	141	A	•	•	•	•	•		1.47	-0.56	•	• .	F	2.12	1.84
Val	142	A	•	•	•	• '	•	•	1.14	-0.99		•	F	1.78	1.77
Thr	143	Ą	•	•	•	•	T	•	0.54	-0.41	•	• ,	\mathbf{F}	1.19	0.55
Gln	144	A:	•	•			Τ.	•	0.54	0.27	•	•	F	0.25	0.31
Asp	145	Α.	•	•	•		T	•	-0.27	0.19	*	•	F	0.25	0.73
Cys	146	A	•	•	:		T		-0.84	0.23	•	:	• .	0.10	0.42
Leu	147	. А	A						-0.58	0.43	•			-0.60	0.17
Gin	148	A	A						-0.27	0.53	. •			-0.60	0.10
Leu	149	Α	Α						-0.57	0.53	٠.	*		-0.30	0.32
Ile	150	Α.	A	•	•	• •	•		-0.57	0.34	٠	•	•	0.30	0.52

Table 9 (continued)

Res P	osition	1.	п	ш	IV ·	v	VI	VII	VIII	IX	x	. X I	X	וונג נו	XIV
Ala	151		Α					C	-0.21	-0.34		•		1.40	0.52
Asp	152	•			• *	T	T		0.39	-0.26		•	F	2.45	0.91
Ser	153			• •		•	T	С	0.08	-0.51		_	. F	3.00	2.00
Glu	154	•	•				T	Ċ	-0.00	-0.71			F	2.70	2.86
Thr	155					• •	T	С	0.89	-0.53	*	•	F	2.40	1.20
Pro	156			•	В			C	1.52	-0.13	•	٠.	F	1.56	1.55
Thr	157				В	T			1.18	-0.51		•	F	1.92	1.79
Ile	158	A			В				1.18	-0.09		•	. F	1.08	1.23
Gln	159					T	T		0.93	-0.19	•	•	F	2.04	1.07
Lys	160					T	T		0.93	0.14	*	• •	F	1.60	1.16
Gly	161					T	Ť		0.44	0.14		•	F	1.44	2.38
Ser	162	•				Ť	T		-0.10	0.14	*	•	F	1.28	1.19
Tyr	. 163				В	·T	·	•	0.58	0.49	*	•	r	0.12	0.44
Thr	164			В	B	-			0.29	0.42	*	•	•	-0.44	
Phe	165 -			В	В		-	•	-0.57	1.40	*	•	•	-0.60	0.69
Val	166			В	В	•	•	. •	-1.03	1.70		٠,	•	-0.60 -0.60	0.54
Pro	167			B	B	•	•	•	-1.03	1.63	.•	• '		-0.60	0.29
Trp	168	A			В	•	•	. •	-1.49	1.53	. •	•	•		0.16
Leu	169 ·	A			В		•	•	-1.13	1.53	*		•	-0.60 -0.60	0.25 0.29
Leu .	170	Α			В			• • •	-0.32	0.89	*	•	. •	-0.30	0.29
Ter	171	A							0.19	0.46	*	•	•	0.20	
Phe	172					T	• 1		0.10	-0.03		•	•	1.80	0.71 0.85
Lys	173 .					T	T	•	-0.20	-0.33		•	F	2.60	1.38
Arg	174	٠.					T	Ċ	±0.20	-0.51			F	3.00	1.04
Gly	175 .						Ť	č	0.61	-0.21	. •	•	F	2.25	0.99
Ser	176	Α					T		0.91	-1.00	•	• .	F	2.05	0.86
Ala	177	Α	• A						1.66	-1.00		•	F	1.35	0.76
Leu	178 ·	Α	$\cdot \mathbf{A}$						1.61	-1.00		•	F	1.20	1.54
Glu	179	A	A						1.50	-1.43		•	F	0.90	1.98
Glu	180	Α	A						1.89	-1.41	*	·	·F	0.90	3.16
Lys	181	Α	A						1.30	-1.91	*		F	0.90	7.66
Glu	182	Α	A			•	. •		1.08	-1.91		•	F	0.90	3.10
Asn	183 🕟	A	A						1.03	-1.23		. •	F	0.90	1.48
Lys	184 🕟	A	A						1.08	-0.59	*		F	0.75	0.55
Πe	185	A	A			•	•		1.08	-0.59	*			0.60	0.63
Leu	186	A	A	• .			•		0.72	-0.59	*	•		0.60	0.68
Val	187	A	A						0.38	-0.50		*	·	0.30	0.49
Lys	188	A	A						0.13	-0.07		. •	F	0.45	0.69
Glu	189	A					T		-0.61	0.00	•	*	F	0.40	1.32
Thr	190		• .		• **	T	T		-0.42	0.10		•	F	0.80	1.54
Gly	191		•	•		T	T	•	-0.50	0.24	•		F	0.65	0.67
Тут	192	•	•	•	•	T	T	•	0.11	0.93	*	•		0.20	0.27
Phe	193	•	•	В	В				-0.28	1.69				-0.60	0.29
Phe	194 .	• `	:	В	В		•		-0.28	1.63	•			-0.60	0.29
Ile	195	• .	•	В	B .			•	-0.82	1.60				-0.60	0.32
Тут	196	•		В	В				-1.29	1.49		:	• .	-0.60	0.28
Gly	197	•	•	• .	В	T			-1.29	` 1.39				-0.20	0.26
Gln	198	•	•	•	В	T			-0.90	1.36	•			-0.20	0.59
Val	199	•	•	•	В	•	•	C	-0.20	1.16				-0.40	0.54
Leu	200	•	. •	•	В	• '	•	C	0.73	0.40	•			-0.10	0.92

Table 9 (continued)

Res Po	sition	I	п	.111	IV	v	VI	VII	VIII.	IX	x	XI	ш	XIII	XIV
						T	T		0.67	-0.03				1.25	1.06
Тут	201	•	•	•	•	T	Ť	•	0.07	0.06	•	•	F	0.80	2.06
Thr	202	•	•	•	•	T		•	0.17		. •	•	F	0.80	3.91
Asp	203	•	•	•	. •	T	T	• .		0.17	•	•	F	1.00	2.52
Lys	204	A	•	•	•	•	T	•	0.43	-0.01	•	•	F		1.73
Thr	205	A	Ą	•	•	•	•	•.	0.90	-0.16	•	•	r	0.60	1.73
Tyr	206	A	A	•	•	•	•	•	1.11	-0.21	•	·	•	0.45	
Ala	207	A	A	•	•	•	•	•	0.61	0.29	•	•	•	-0.30	0.70
Met	208	Α	A	•	<u>.</u>	•	•	•	-0.28	0.97	:	•	•	-0.60	0.40
Gly	209	A	A		В.	•	•	. •	-0.32	1.17	-	•	•	-0.60	0.18
His	210	A	A	•	В	•	•	•	0.10	0.81	•	·• ·	•	-0.60	0.31
Leu	211	A	\mathcal{A}	•	В	•	•	•	0.39	0.31	•	•	•	-0.30	0.61
Ile	212	A	A	•	В		•	•	1.02	-0.30	•	:	•	0.45	1.22
Gln	213	A	. A		В	•	•	•	0.77	-0.73	•		•	0.75	1.80
Arg	214	Α	A	•	В	. •	•	•	1.08	-0.59	•	•	F	0.90	1.62
Lys	215	Α	A	•	. В	•	•	• .	0.26	-0.77	*		F	0.90	3.14
Lys	216	Α	Α		В	•	• *	•	0.37	-0.81			F	0.90	1.35
Val	217		Α	В	В		•	•	0.91	-0.43	*	*	•	0.30	0.60
His	218		A	В	В		•		0.91	-0.00		*	.	0.30	0.29
Val	219		A	В	В				. 0.80	-0.00	*	*	•	0.30	0.25
Phe	220		•	В	В				-0.06	-0.00	*	:		0.30	0.57
Gly	221	Α			В				-0.40	0.04		*	•.	-0.30	0.35
Asp	222	Α				• .		•	-0.36	-0.07	*	: .	•	0.50	0.63
Glu	223	· A						• .	-1.18	-0.03	*		.•	0.50	0.60
Leu	224	Α			В			•	0.63	-0.17		• '	•	0.30	0.45
Ser	225.	. A			В				-0.74	-0.11				0.30	0.39
Leu	226	· A			В		• .		-1.10	0.57		•	•	-0.60	0.18
Val	227	· A			В				-0.99	1.36	•	*		-0.60	0.19
Thr	228	Α			В		•	. •	-1.66	0.67		*		-0.60	0.28
Leu		A			В				-1.73	0.86	•			-0.60	0.18
Phe		· · A		•	В				-1.43	0.86.		-		-0.60	0.17
Arg	231	· A			. B				-0.62	0.61	*			-0.60	0.21
Cys	232		-		В.	T			-0.37	0.53				-0.20	0.41
Ile	233				В	T			-0.27	0.46				-0.20	0.46
Gln	234				В	T			0.54	0.10	*			0.10	0.37
Asn	235				В			С	0.93	0.10	*			0.05	1.19
Met	236	,]	·		В			С	0.01	0.01			F	0.20	2.44
Pro	237	•			В			C	0.47	0.01			F	0.44	1.16
Glu	238	•	Ť	·		T	•		1.36	0.04	•		F	1.08	1.12
Thr	239	•	•	•				C	1.36	0.04	*		F	1.12	1.82
Leu	240	•	•	•				C	1.06	-0.17			F	1.96	1.89
Pro	241	•	•	•		T	-		0.99	-0.21			F	2.40	1.46
Asn	242	•	•	•	•	Ť			0.96	0.36			F	1.41	0.54
Asn	243	•	•	•	•	Ť	T		0.66	0.63			F	1.22	1.03
Ser	244	•	•	•	. •	Ť	Ť		0.38	0.33			F	1.13	0.89
	245	•	:	•	•	Ť	Ť	•	0.84	0.40	-		-	0.74	0.56
Cys		•	•	•	•	Ť	Ť	:	0.17	0.43	•	•	·	0.20	0.35
Tyr	246		•	•	•	. •		•	-0.42	0.71	•	•	•	-0.40	0.18
Ser	247	A A		•	•	•	•	•	-0.38	0.83	•	•	. •	-0.60	0.34
Ala	248		A	•	• •	•	•	•	-0.89	0.26	•	•	.•	-0.30	0.43
Gly	249	A	A	•	•	•	•	•	-0.22	0.19		•	•	-0.30	0.43
Ile	250	A	A			•	•	•	-0.22	0.13		•	•	- 0.50	U.21

Table 0	(continued)	
IXINETI	e communicary	

Res Po	sition	1.	11	щ	IV ·	v	VI	VII	VIII	IX	x .	IX	XII	XIII	XIV
Ala	251	. A	A					•	0.02	-0.20	*	•		0.30	0.46
Lys	252	Α	A.						-0.02	-0.70	•			0.60	0.80
Leu	253	A	A	•			• .	•	0.57	-0.70			F	0.90	1.13
Glu	254	A.	A				•		0.91	-1.39		. •	F	0.90	1.87
Glu	255	Α	Α		•	•:			0.99	-1.89			F	0.90	1.62
Gly	256	A	A:	•	•	•	٠.		1.58	-1.20		• •	F	0.90	1.62
Asp	257	Α.	A	•	•				0.72	-1.49		*	F	0.90	1.62
Glu	258	. A	A				•	•	0.94	-0.80	*	*	F	0.75	0.77
Leu	259	A	. A .					•	0.06	-0.30	*	•		0.30 -	0.79
Gln	260	Α	A	•.	•			• .	-0.16	-0.04	*	•		0.30	0.33
Leu	261	Α	A	•			• .	•	0.30	0.39	*			-0.30	0.30
Ala	262	. A	A	•					0.30	0.39	*			-0.30	0.70
Ile	263	A	Α	•	•			• •	0.30	-0.30		•		0.30	0.70
Pro	264	Α	•				T		0.52	-0.30		*	F	1.00	1.37
Arg	265	Α					T	•	0.52	-0.49		*	F	1.00	1.37
Glu	266	Α					T	•	0.44	-0.59	. *	•	F	1.30	3.38
Asn	267	A	•	. •	•		T	•	0.73	-0.59		*	F	1.30	1.53
Ala	268	A.				•		•	0.81	-0.63	* .	*	:	0.95	1.05
Gln	269	• A		•	. •		•		1.02	0.06	*	•.		-0.10	0.50
Ile .	270	Α	• •	•		•	•	•	0.57	0.06		• '		0.15	0:52
Ser	. 271				•. •		•	С	0.57	0.09		*		0.60	0.51
Leu	272	٠.					• 4	C	-0.29	-0.41		*	F	1.60	0.49
Asp	273			٠.	•	T	_ T	•	-0.01	-0.17		•	F	2.25	0.52
Gly	274		•		•	T	T	•	-0.71	-0.37		*	F	2.50	0.56
Asp	275	•		•	•	T	T		-0.52	0.03		•	F	1.65	0.59
Val	276	A			•	•	T	•	-0.57	0.13		*	F	1.00	0.30
Thr	277	A	•		В		• .		-0.34	0.56	•	*		-0.10	0.30
Phe	278	A			В	•		•	-1.16	0.63		•		-0.35	0.18
Phe	279	· А			В	•	•	• '	-0.77	1.31		• .		-0.60	0.20
Gly	280	. A	Α						-1.58	0.67		. ●	٠.	-0.60	0.28
Ala	281	A	A			•	•		-1.53	0.87		• .		-0.60	0.27
Leu	282	Α	Α				. •	•	-1.61	0.77	•	•		-0.60	0.26
Lys	283	A	A	• '	• •	•	٠.		-1.30	0.41	*	•		-0.60	0.33
Leu	284	· A	A				•		-0.99	0.41			•.	-0.60	0.42
Leu	285	A	A	•		. •	•	•	-1.03	0.34	*		•	-0.30	0.65

Ta	L	•	1	Ω
14	D1	c		v

Res F	Position	1	. п	. III	IV	V	· VI	VII	VIII	ΙX	X	. XI	XII	XIII	XIV
Met	. 1	A							0.73	-0.71				0.95	1.39
Asp	2	A					T		1.12	-0.66		•	•	1.15	1.56
Asp	. 3	Α					T		1.62	-1.09	* .	•	•	1.15	2.12
Ser	4	Α					Ť	٠,	2.01	-1.51		•	•	1.15	4.19
Thr	5	Ā			• .	•	Ť	•	2.40	-2.13	•	•	F	1.13	-
Glu	6	Ā	A	·	·	•	•	•	2.70	-1.73			F		4.35
Arg	7	A	Ā	•	•	•	•	•	2.81	-1.73		•		0.90	4.51
Glu	8	A	A	•	•	•	•	•	2.00			Ţ	F	0.90	4.51
Gln	Š	A	- A	. •	•	•	•	•		-1.73		- T	F	0.90	6.12
Ser	10	Ā	•	•	В	•	•	. •	1.99 2.00	-1.53			F	0.90	2.91
Arg	11	Ā	•	. •	B.	•	•	•		-1.04	Ŀ	·•	F	0.90	2.15
Leu	12	Â	•	•	В	•	•	•	1.33	-0.66	I		F	0.90	1.66
Thr	13	Ā	•	•	В	•	•	•	0.41	-0.09	•		F	0.45	0.51
Ser-	14	A	A	•	.0	•	. •	•	0.46	0.20			F	-0.15	0.32
	15	A		• '	•	• •	•	•	0.50	-0.19	*		•	0.30	0.32
Cys Leu	16		A	•	•	•	•	• .	0.91	-0.19	*		•	0.30	0.78
		Α.	A	•	•	• .	• '	• .	0.80	-0.87	*	*.	F	0.90	1.06
Lys	17	A	. A	•	• • .	•	•	•	1.61	-1.36	•	•	F	0.90	1.37
Lys	18	A	A	•	•	•	• .	•	1.32	-1.74	• .	*	F.	0.90	4.44
Arg	19	A	A	•	. •	•	•	•	1.67	-1.70		* '	F	0.90	5.33
Glu	20	A	Α	• .	•		•	•	1.52	-2.39	•	•	F	0.90	5.33
Glu	21	Ą	A	•	• .	• .	•	•	2.38	-1.70		*	F	0.90	2.20
Met	22	A	Ą	•	•	• .	•	•	2.33	-1.70		: • •	F	0.90	2.24
Lys	23	A	A	•	· •	•	•	• .	1.62	-1.70	*	*	F	0.90	2.24
Leu	24	A	A	•	•	• '			0.66	-1.13		*	F	0.75	0.69
Lys	25	A	· A		•	•			0.36	-0.49		•	F.	0.45	0.52
Glu	26	A	A	•	В	•	• .		-0.53	-0.71	*			0.60	0.35
Cys	27	A	Α	•	В				-0.74	-0.03	*			0.30	0.30
Val	28	· A.	A		В	. •	• .	. •	-1.00	-0.03	•	*		0.30	0.12
Ser	29	Α	Α.	. •	В				-0.08	0.40	•	• 1		-0.30	0.11
Ile	30	Α			В				-0.08	0.40	*	•		-0.30	0.40
Leu	31	Α			В				-0.08	-0.17	*			0.45	1.08
Pro	32				В.			С	0.29	-0.81	•		F	1.10	1.39
Arg	33					T			0.93	-0.81			F	1.50	2.66
Lys	. 34		• •		•	T			0.93	-1.07			F	1.84	4.98
Glu	35				•			C	0.97	-1.37	* -	•	F	1.98	4.32
Ser	36						T	С	1.89	-1.16	•		F	2.52	1.64
Pro	37						T ·	Ċ	1.80	-1.16		*	F	2.86	1.60
Ser	38	•				T	T		1.39		*		F	3.40	1.24
Val	. 39	A					Ť	•	1.39	-0.39			F	2.36	
Arg	40	. · A					-		1.39	-0.77	•		F	2.46	1.24
Ser	41	· A			·	•	•	•	1.34	-1.20	•		F	2.46	
Ser	42			·	·	T.	T ·	•	1.60	-1.16		•	r		2.00
Lys	43		•	•	•	Ť	Ť	•	1.09	-1.10				3.06	2.67
Asp	44	•	•	• .	•	Ť	Ť	• .	1.13		•	Ĭ	F	3.06	2.72
Gly	45	Å	•	• .	•	•	Ť	•		-1.11	*	I	F	3.40	1.67
Lys	46	A	A	•	•	•		•	0.43	-0.81	~	•	F	2.66	1.03
Leu	47	Â	A	•	•	•	•	•	0.14	-0.70		•	F	1.77	0.52
Leu	48	A	A	•	•	•	•	•	0.13	-0.20	. .	•	•	0.98	0.31
Ala	49	A	A	•	• .	•	•	•	-0.72	0.29	-	:	. •	0.04	0.46
Ala	50			•	•	• .	:	•	-1.53	0.54	•	-	•	-0.60	0.19
UIR	20	$\mathbf{A}_{\mathbf{c}}$, A	•	• ·	•	•	•	-2.00	1.23	• .	•	•	-0.60	0.19

Table 10 (continued)

Res Po	sition	I	. п	m	IV	v	. VI	VII.	VIII .	IX	x	XI	XII	XIII	XIV
Thr	51	· A	Α				•	•	-2.63	1.23	•	٠.	•	-0.60	0.19
Leu	52	A	A	٠.			•	•	-2.63	1.04		•	•	-0.60	0.19
Leu	- 53	. A	À					•	-2.63	1.23	•	•	•	-0.60	0.15
Leu	54	A	Α		•		•	• 1	-2.34	1.41	. •	•	•	-0.60	0.09
Ala	55	Α	Α		••		•	• .	-2.42	1.31	•	• .		-0.60	0.14
Leu	56	A	Α			•	•	•	-2.78	1.20	. •	• .	• *	-0.60	0.09
Leu	57	A			•	•	T	•	-2.78	1.09	•	. •	•	-0.20	0.06
Ser	58	Α			•	•	T		-2.28	1.09	•	•	. •	-0.20	0.05
Cys	59	A	•		•	•	T	.•	-2.32	1.07	٠.	•	•	-0.20	0.09
Cys	60	· A		•	•	•	T	•	-2.59	1.03	٠.	• .	•	-0.20	0.08
Leu -	61		. •	В	В	•	•	•	-2.08	0.99	•.	.•	•	-0.60	0.04
Thr	62		•	В	В			•	-1.97	0.99	•	•	•	-0.60	0.11
Val	63			В	В	•	•	•	-1.91	1.20	•	•	•	-0.60	0.17
Val	64		•	В -	В		•	•	-1.24	1.39	.•	•	• -	-0.60	0.33
Ser	65	•	•	В	. B	•	•	• .	-1.43	1.10	•	•	•	-0.60	0.40
Phe	66	A	•	•	В	•	•	•	-1.21	1.26	. •	•• .	•	-0.60	0.40
Tyr	67	A	•	•	В.	•	•	•	-1.49	1.11	•	•	• .	-0.60	0.54
Gin	68	A	•	•	В	. •	• '	•	-1.44	0.97	• .	•	• .	-0.60	0.41
Val	69	A	•	•	В	•	•	•	0.59	1.27	•	•	• .	-0.60	0.39
Ala	70	Α	•		B	•	. •	•	-0.63	0.89	•	•	. •	-0.60	0.43
Ala	.71	A		•	В	• .	· <u>·</u>	•	0.07	0.56	•	. •	•	-0.60	0.25
Leu	72	A	•	•	•	• .	T	•	-0.50	0.16	•	•		0.10 0.25	0.55
Gln	73	A	•	•		•	T	• .	-1.09	0.20	• .	• .	F F	0.25	0.45 0.45
Gly	74	A	•	•	•	•	T	•	-0.53	0.20 0.09	•		F	0.25	0.43
Asp	75	, A	•	•	•	• .	T	•	-0.76 -0.06	0.09	•		F	-0.15	0.75
Leu	76	A	Ą	•	•	•	•	. •	-0.06 0.17	-0.31	٠.		r	0.30	0.55
Ala	<i>7</i> 7	A	Ą	٠	•	•	• •	•	0.17	-0.24	•		•	0.30	0.42
Ser	78	A	A	•	•	. •	•	• •.	-0.30	-0.24	•		•	0.30	0.42
Leu	79	A	A	•	•		•	•	-0.30	-0.24	•	*	• .	0.30	0.72
Arg	80	A	A	• •	•	•	•	•	0.17	-0.34	•	•	. •	0.30	0.93
Ala	81 82	A A	A A	•	•	•	•	•	0.72	-0.30	•	•	•	0.45	1.11
Glu	82 83	A	A	. •	• .	•	•	•	0.72	-0.49	•		•	0.30	0.77
Leu Gln	· 84	A	A	•	• .	•	•	• .	1.21	0.01	•		•	-0.15	1.04
Gly ·	85	A	. A	. •	•	•	•	• 0	1.10	0.01			·	-0.30	0.61
His	86	·A	Ā	•	•	. •	•	•	1.73	0.01	*			-0.15	1.27
His	87	Ā	Â	•	•	•		•	0.92	-0.67			-	0.75	1.47
Ala	88	i A	Ā	•	•	•	•	•	1.52	-0.39	Ċ	• 1		0.45	1.22
Glu	89	A	A	•	•	•	. j	•	0.93	-0.39				0.45	1.39
Lys	90	· A	A	•	•	•		:	0.93	-0.39	•		F	0.60	1.03
Leu	91	A	• •	•	·		T		0.38	-0.46	*	÷		0.85	1.01
Pro	92	• A		•			T		0.07	-0.46				0.70	0:59
Ala	93	Ä	•	•			Ť		0.07	-0.03			•	0.70	0.29
Gly	94	Ā	•				T		-0.14	0.47				-0.20	0.36
Ala	95	A	:						-0.14	0.21		*		-0.10	0.36
Gly	96	A		•					0.08	-0.21	•		F	0.65	0.71
Ala	97	A				-			-0.06	-0.21			F	0.65	0.72
Pro	98	A							-0.28	-0.21		•	F	0.65	0.71
Lys	99	A	A	•					0.07	-0.03		•	F	0.45	0.59
Ala	100	A	Ā		• .				0.66	-0.46			F	0.60	1.01

Table 10 (continued)

Res Po	sition	I.	п	ш	IV -	v .	VI	VII	VIII	IX	x	ж	ш	XIII	XIV
Gly	101	. А	A						0.41	-0.96			F	0.90	1.13
Leu	102	Â	Â	•	•	•	•	•	0.79	-0.89	·	•	F	0.75	0.57
	102	A	Â	•	•	•	•	•	0.79	-0.46	•	•	F	0.45	0.88
Glu				• '	•	•	•	.*			•	•	F	0.45	0.89
Glu	104	A	A	•	•	•	•	•	-0.49	-0.46 -0.24	-	. •	r	0.43	0.81
Ala	105	A	A	•	•	•	•	•	-0.21		•	•	•	0.30	_
Pro	106	A	A	•	•	•	•	•	-0.46	-0.44	•	•	• .		0.67
Ala	107	A .	A	•	•	•	. •	•	0.01	0.06	:	:	•	-0.30	0.39
Val	108	A	A	•	•	•	•	•	-0.80	0.49	•		•	-0.60	0.38
Thr	109	Ą	Ą	•	•	•	•	•	-0.76	0.67	:	- I	•	-0.60	0.20
Ala	110	Ą	A	•.	•	•	•	• .	-1.06	0.24	Ţ.	-	•	-0.30	0.40
Gly	111	, A	A	•	•	•	• .	• .	-1.54	0.43	•	•	•	-0.60	0.38
Leu	112	Α	A	. •	•	•	•	•	-0.96	0.57	*	*	•	-0.60	0.23
Lys	113	•	A	В	•	. •	•	•	-0.31	0.09		•	•	-0.30	0.39
Ile	114	•	A	В	•	•	•	•	-0.21	0.01	•	•	•	-0.30	0.61
Phe	115	•	Α	В	•	•	•	•_	-0.21	0.01	*	•	•	0.15	1.15
Glu	116	•	A	•	•	•	•	C	-0.08	-0.17	*	•	F	1.25	0.58
Pro	117	•	A	. •	•	•	•	С	0.39	0.26	. *	*	F	1.10	1.28
Pro	118	•	•	•	•	•		C	0.34	0.00	*		F	2.20	1.47
Ala	119	•	•		•	•	T	С	0.89	-0.79		• .	F	3.00	1.47
Pro	120		• •	•			T	С	1.59	-0.36		*	F	2.25	0.94
Gly	. 121			•		Τ.	T		1.29	-0.39	• *	*	\mathbf{F} .	2.15	0.98
Glu	122			•		T	T	•.	1.20	-0.43	•		F	2.00	1.30
Gly	123				•			Ċ	1.41	-0.54		•	F	1.60	1.12
Asn	124						T	С	2.00	-0.57			F	1.50	1.97
Ser	125			•			T	С	1.91	-0.60		•	F	1.50	1.82
Ser	126			•			T	С	2.37	-0.21		*	F	1.54	2.47
Gln	127						T	С	2.37	-0.64		*	F	2.18	3.01
Asn	128							С	2.76	-0.64			F	2.32	3.61
Ser	129	٠.			:		T	С	2.87	-1.03			F	2.86	5.39
Arg	130					T	T		2.58	-1.41			F	3.40	6.09
Asn	131					T	T		2.02	-1.31	*		F	3.06	3.83
Lys	132					Т	T		2.02	-1.07	*		F	2.72	2.12
Arg	133					T			1.68	-1.06			F	2.18	1.88
Ala	134							. C	1.77	-0.63	•		F	1.64	1.15
Val	135							С	1.66	-0.60	*		F	1.15	0.89
Gln	136							С	1.66	-0.60	*		F	1.49	0.79
Gly	137	-					T	С	1.30	-0.60	*		F	2.18	1.35
Pro	138	-					T	С	0.84	-0.61	*		F	2.52	2.63
Glu	139			_			T	С	1.13	-0.83	•		F	2.86	1.50
Glu	140		• .			T	T		1.74	-0.84			F	3.40	2.03
Thr	141					Ť			1.43	-0.51			F	2.86	2.06
Gly	142	•	•	•	•	Ť	T		1.08	-0.46		. *	F	2.42	1.72
Ser	143	•	•	•	•	Ť	Ť	·	0.43	0.33			F	1.33	0.86
Туг	144	•,	•	•	•	Ť	Ť	•	0.22	0.97	·.	• .	-	0.54	0.44
Thr	145	•	•	•	•	Ť	Ť	•.	-0.07	0.91	•	•	•	0.20	0.69
Phe	146	• .	•	В	В	•	-	•	-0.57	1.40	•	•	•	-0.60	0.54
Val	147	•	•	В	В	•	•	•	-1.03	1.70		•	•	-0.60	0.29
Pro	148	•	•	В	В	•	•	•	-1.03	1.63		• .	•	-0.60	0.16
	148	A	•		В	•	•		-1.49	1.53	٠.	•	•	-0.60	0.15
Trp		A	•	•	В	• .	•	•	-1.13	1.53	•		•	-0.60	0.23
Leu	150	A.	•	•	D	•	•	•	-1.13	1.33	-	•	•	-0.00	U.L.7

Table 10 (continued)

Res P	osition	I	. 11	,m	IV	v	. VI	VII	VIII	1X	x	XI	XII	XIII	XIV
Leu	151	A		•	В				-0.32	0.89	*			-0.30	0.38
Ser	152	Α							0.19	0.46	*	•	•	0.20	0.71
Phe	153		•			T			0.10	-0.03	** *	•	•	1.80	0.85
Lys	154					T	T		-0.20	-0.33	*	•	F	2.60	1.38
Arg	155					-	T	Ċ	-0.20	-0.51		•	F	3.00	1.04
Gly	156						Ť	č	0.61	-0.21	•	•	F	2.25	
Ser	157	A			·	·	Ť	•	0.91	-1.00	•	•	F	2.25	0.99
Ala	158	A	A				•	•	1.66	-1.00		. •	F	1.35	0.86
Leu	159	A	Ā	·		·	•	•	1.61	-1.00		. •	F		0.76
Glu	160	Ā	Ā		·	•	• .	. •	1.50	-1.43	•	•	F	1.20	1.54
Glu	161	A	A	•	· • .	•	•	•	1.89	-1.41		• .		0.90	1.98
Lys	162	A	Ā	•	•	•	•	•	1.30	-1.91	•	•	F	0.90	3.16
Glu	163	Ā	Ā	•	•	.•	•	•	1.08	-1.91		• .	F	0.90	7.66
Asn	164	Ā	Ā	•	•	•.	•	•	1.03	-1.23	*		r F	0.90	3.10
Lys	165	Ā	Ā	•	•	•	•	•	1.08	-0.59	•	-	_	0.90	1.48
Ile	166	Ä	Ä	•	. •	•	•	• .	1.08	-0.59		:	F	0.75	0.55
Leu	167	· A	A	•	•	•	•	• '	0.72	-0.59 -0.59			•	0.60	0.63
Val	168	Ā	Ā	•	•	•	•	•	0.72	-0.50	•		•.	0.76	0.68
Lys	169	A	A	•	•	•	• .	•	0.13	-0.07			F	0.92	0.49
Glu	170	A		•	•	•	T	•	-0.61	0.00	•		F	0.93	0.69
Thr	171	••	•	•	•	T.	T	•	-0.42	0.10		¥	F	1.64	1.32
Gly	172	•		·	•	Ť	Ť	•	-0.50	0.10	•		F	1.60 1.29	1.54
Tyr	173			•	·	Ť	Î	•	0.11	0.93	•		r	0.68	0.67
Phe	174			В	В	-	•	• .	-0.28	1.69			: .	-0.28	0.27
Phe	175			В	B		•	•	-0.28	1.63	•	•	•	-0.26 -0.44	0.29 0.29
Ile	176			В	В				-0.82	1.60	•		• .	-0.60	0.29
Tyr	177			В	B				-1.29	1.49	•	· ·	•	-0.60	0.32
Gly	178	٠.			В	T			-1.29	1.39	•	•	•	-0.20	0.26
Gln	179				В	T			-0.90	1.36	•	•	•	-0.20	0.20
Val	180	٠.			В		·	C	-0.20	1.16	•	•	•	-0.40	0.54
Leu	181				В			Č	0.73	0.40		•	•	-0.10	0.92
Tyr	182					T	T		0.67	-0.03	•	• .,	•	1.25	1.06
Thr	183					T	T		0.77	0.06		•	F	0.80	2.06
Asp -	184					T	T		0.18	0.17		•	F	0.80	3.91
Lys	185	A					T		0.43	-0.01		•	F	1.00	2.52
Thr	186	Α	A		•				0.90	-0.16		•	F	0.60	1.73
Tyr	187	Α	A						1.11	-0.21	•	•		0.45	1.03
Ala	188	Α	. A					•	0.61	0.29		•	•	-0.30	0.70
Met	. 189	A	A						-0.28	0.97				-0.60	0.40
Gly	190	. A	A		В			٠.	-0.32	1.17		-		-0.60	0.18
His	191	Α	· A		В		:		0.10	0.81	*			-0.60	0.31
Leu ·	192	Α	A		В.				0.39	0.31				-0.30	0.61
Ile	193	A	Α		В				1.02	-0.30			-	0.45	1.22
Gln	194	A	Α		В			. •	0.77	-0.73		•	·	0.75	1.80
Arg	195	A	A ·		В			·.	1.08		*		F	0.90	1.62
Lys	196	A	Α		В				0.26	-0.77	•	*	F	0.90	3.14
Lys	197 .	A	A		В				0.37	-0.81		*	_	0.90	1.35
Val	198		A	В	B				0.91	-0.43	• •	•		0.30	0.60
His	199		Α	В	В				0.91	0.00	*	•	•	0.30	0.29
Val	200		Α	В	B.	•	•	•	0.80	0.00	*	•		0.30	0.25

Table 10 (continued)

	-														
Res Po	sition	I.	п	Ш	IV -	Ÿ.	VI	VII	VIII	IX	x	XI	XII	XIII	VIX
Phe	201			В	В				-0.06	0.00	•	•		0.30	0.57
Gly	202	Α			В				-0.40	0.04	:	*		-0.30	0.35
Asp	203	A		• •					-0.36	-0.07	•			0.50	0.63
Glu	204	A						:	-1.18	-0.03	*		•	0.50	0.60
Leu	205	Α			В	• .	•		-0.63	-0.17			•	0.30	0.45
Ser	206	Α		•	В	•		•	-0.74	-0.11	•	•		0.30	0.39
Leu	207	A			В				-1.10	0.57		•		-0.60	0.18
Val	208	Α			В		•		-0.99	1.36	•	•	. •	-0.60	0.19
Thr	209	Α			В	•			-1.66	0.67	*	•	• '	-0.60	0.28
Leu	210	A			В				-1.73	0.86		• .		-0.60	0.18
Phe	211	Α		•	В		• .	•	-1.43	0.86	*	•		-0.60	0.17
Arg	212	A		·.	. B		•	•	-0.62	0.61	•	٠.	•	-0.60	0.21
Cys	213			•	В	, T		•	-0.37	0.53	*	•	•	-0.20	0.41
Ile	214				B	T		. •	-0.27	0.46	*		•	-0.20	0.46
Gln	215	•		•	В	T		. •	0.54	0.10	*	•	•	0.10	0.37
Asn	216				В			· C	0.93	0.10	*		• *	0.05	1.19
Met	217		•		· B		•	С	0.01	0.01	*		F	0.20	2.44
Pro	218				В	•		. C	0.47	0.01	•		F	0.44	1.16
Glu	219		• •	•	. •	T		• .	1.36	0.04	*		F	1.08	1.12
Thr	220				•			С	1.36	0.04	*	•	F	1.12	1.82
Leu	. 221				• .	•	•	С	. 1.06	-0.17	*.	•	F .	1.96	1.89
Pro	222					T	• ,	•	0.99	-0.21·	•	•	F	2.40	1.46
Asn	223	•	•		•	T	•	• •	0.96	0.36	•	•	F ·	1.41	0.54
Asn	224		•	•		T	T	•	0.66	0.63		•	F	1.22	1.03
Ser	225			•	•	T	T	•	0.38	0.33	•	•	F	1.13	0.89
Cys	226			•		T	T	-	0.84	0.40	•	•	. •	0.74	0.56
Tyr	227				•	T	T		0.17	0.43	•	•		0.20	0.35
Ser	228	Α	• •		•	• .	• '		-0.42	0.71	. •	•	•	-0.40	0.18
Ala	229	Α	A				•	•	-0.38	0.83	•		•	-0.60	0.34
Gly	230	· A	A		•	•	•	•	-0.89	0.26	•	. •	· •	-0.30	0.43
Ile	231	Α	Α		•	•	•	•	-0.22	0.19	*	• .	•	-0.30	0.27
Ala	232	A	A		•	•	. •	•	0.02	-0.20	•	•	•	0.30	0.46
Lys	233	· A	A	•	•	•	•	•	-0.02	-0.70	•	• '	<u>:</u>	0.60	0.80
Leu	234	A	Α	•	•	•	• .	• •	0.57	-0.70	•	•	F	0.90	1.13
Glu	235	Α	A	•	•	•	•	•	0.91	-1.39	•	•	F	0.90	1.87
Glu	236	Α	A	• .	•	. •	•	•	0.99	-1.89	•	:	F	0.90	1.62
Gly	237	Α	Α	•	•	. •	•	•	1.58	-1.20	•		F	0.90	1.62
Asp	238	· A	A	•	•	•	•	•	0.72	-1.49			F	0.90	1.62
Glu	239	A	Α	. •	•	•	•	•	0.94	-0.80	•		F	0.75	0.77
Leu	240	Α.	A	. •	• **	•	•	•	0.06	-0.30		. •	•	·0.30	0.79
Gln	241	Ą	A	•	. •	•	•	•	-0.16	-0.04		٠.	•	0.30	0.33
Leu	242	A	A	•	•	•	•	•	0.30	0.39	Ţ	•	•	-0.30	0.30
Ala.	243	A.	Α	•	•	•	•	•	0.30	0.39	Τ.		• •	-0.30	0.70
Ne	244	A	Α	•	•	•	<u>.</u>	•	0.30	-0.30	•	•		0.30	0.70
Pro	245	Ą	. •	•	• .	•	T	• .	0.52	-0.30	•	:	F	1.00	1.37
Arg	246	A.	•	•	•	•	T	•	0.52	-0.49	:	-	F	1.00	1.37
Glu	247	A	•	•	•	•	T	•	0.44	-0.59		Ξ	.F	1.30	3.38
Asn	248	Ą	•	•	•	•	T	•	0.73	-0.59	-		ŗ	1.30	1.53
Ala	249	A	•	•	•	•	•	•	0.81	-0.63	-		•	0.95	1.05
Gln	250	A	•	•	•	• .	•	•	1.02	0.06	•	-	•	-0.10	0.50

Table 10 (continued)

Res Po	sition	1.	П	ш	. VI	V	VI	VII	vIII	IX	x	IX	ХП	XIII	XIV
Ile	251	. A	•			·			0.57	0.06	*	•		0.15	0.52
Ser	252							С	0.57	0.09	•	*		0.60	0.51
Leu	253							·C	-0.29	-0.41		• .	F	1.60	0.49
Asp	254			•		T	T		-0.01	-0.17		*	F	2.25	0.52
Gly	255					T	T		-0.71	-0.37		. *	·F	2.50	0.56
Asp	256			•		T	T		-0.52	0.03		•	F.	1.65	0.59
Val	257	A					T	•	-0.57	0.13		*	F	1.00	0.30
Thr	258	A			В				-0.34	0.56		*		-0.10	0.30
Phe	259	A			В	٠.			-1.16	0.63		*	•	-0.35	0.18
Phe	260	A			В				-0.77	1.31	• .	• 1		-0.60	0.20
Gly	261	. A	A		٠.				-1.58	0.67			•	-0.60	0.28
Ala	262	A	Α.			•	•		-1.53	0.87				-0.60	0.27
Leu	263	A	A	•		. •			-1.61	0.77	*			-0.60	0.26
Lys	264	A	A		•		· .		-1.30	0.41	•	•		-0.60	0.33
Leu	· 265	A	A	•					-0.99	0.41				-0.60	0.42
Leu	266	A	A		•			•	-1.03	0.34	•		• .	-0.30	0.65

[083] In another embodiment, the invention provides antibodies that bind a polypeptide comprising, or alternatively consisting of, an epitope-bearing portion of a polypeptide of the invention. The epitope of this polypeptide portion may be an immunogenic or antigenic epitope of a polypeptide of the invention. An "immunogenic epitope" is defined as a part of a protein that elicits an antibody response when the whole protein is the immunogen. On the other hand, a region of a protein molecule to which an antibody can bind is defined as an "antigenic epitope." The number of immunogenic epitopes of a protein generally is less than the number of antigenic epitopes. See, for instance, Geysen et al., Proc. Natl. Acad. Sci. USA 81:3998-4002 (1983).

[084] As to the selection of polypeptides bearing an antigenic epitope (i.e., that contain a region of a protein molecule to which an antibody can bind), it is well known in that art that relatively short synthetic peptides that mimic part of a protein sequence are routinely capable of eliciting an antiserum that reacts with the partially mimicked protein. See, for instance, Sutcliffe, J. G., Shinnick, T. M., Green, N. and Learner, R. A. (1983) "Antibodies that react with predetermined sites on proteins", Science, 219:660-666. Peptides capable of eliciting protein-reactive sera are frequently represented in the primary sequence of a protein, can be characterized by a set of simple chemical rules, and are confined neither to immunodominant regions of intact proteins (i.e., immunogenic epitopes) nor to the amino or carboxyl terminals. Antigenic epitope-bearing peptides and polypeptides of the invention are therefore useful to raise antibodies, including monoclonal antibodies, that bind specifically to a polypeptide of the invention. See, for instance, Wilson et al., Cell 37:767-778 (1984) at 777.

[085] In specific embodiments, antibodies of the present invention bind antigenic epitope-bearing peptides and polypeptides of BLyS and preferably contain a sequence of at least 4, at least 5, at least 6, at least 7, more preferably at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 20, at least 25, at least 30, at least 40, at least 50, and, most preferably, between about 15 to about 30 amino acids contained within the amino acid sequence of a BLyS polypeptide. Preferred polypeptides comprising immunogenic or antigenic epitopes are at least 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, or 100 amino acid residues in length. Additional non-exclusive preferred antigenic epitopes include the antigenic epitopes disclosed herein, as well as portions thereof.

Non-limiting examples of antigenic polypeptides or peptides that can be [086] used to generate BLyS-specific antibodies and which may be bound by the antibodies of the invention include: a polypeptide comprising, or alternatively consisting of, amino acid residues from about Phe-115 to about Leu-147 in SEQ ID NO:3228; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Ile-150 to about Tyr-163 in SEO ID NO:3228; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Ser-171 to about Phe-194 in SEO ID NO:3228; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Glu-223 to about Tyr-246 in SEQ ID NO:3228; and a polypeptide comprising, or alternatively consisting of, amino acid residues from about Ser-271 to about Phe-278 in Figures 1A and 1B (SEO ID NO:3228). In this context, "about" means the particularly recited ranges and ranges larger or smaller by several, a few, 5, 4, 3, 2 or 1 amino acid residues at either or both the amino- and carboxy-termini. These polypeptide fragments have been determined to bear antigenic epitopes of the BLyS polypeptide by the analysis of the Jameson-Wolf antigenic index, as disclosed Table 9, above.

[087] Non-limiting examples of antigenic polypeptides or peptides that can be used to generate BLyS-specific antibodies and which may be bound by the antibodies of the invention include: a polypeptide comprising, or alternatively consisting of, amino acid residues from about Pro-32 to about Leu-47 in SEQ ID NO:3229; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Glu-116 to about Ser-143 in SEQ ID NO:3229; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Phe-153 to about Tyr-173 in SEQ ID NO:3229; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Pro-218 to about Tyr-227 in SEO ID NO:3229; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Ala-232 to about Gln-241 in SEQ ID NO:3229; a polypeptide comprising, or alternatively consisting of, amino acid residues from about Ile-244 to about Ala-249 in SEQ ID NO:3229; and a polypeptide comprising, or alternatively consisting of, amino acid residues from about Ser-252 to about Val-257 in SEQ ID NO:3229. In this context, "about" means the particularly recited ranges and ranges larger or smaller by several, a few, 5, 4, 3, 2 or 1 amino acid residues at either or both the amino- and carboxy-termini. These polypeptide fragments have been determined to bear antigenic epitopes of the BLyS polypeptide by the analysis of the Jameson-Wolf

antigenic index, as disclosed in Table 10 generated by the Protean component of the DNA*STAR computer program (as set forth above).

[088] BLyS epitope-bearing peptides and polypeptides may be produced by any conventional means. See, e.g., Houghten, R. A. (1985) General method for the rapid solid-phase synthesis of large numbers of peptides: specificity of antigen-antibody interaction at the level of individual amino acids. Proc. Natl. Acad. Sci. USA 82:5131-5135; this "Simultaneous Multiple Peptide Synthesis (SMPS)" process is further described in U. S. Patent No. 4,631,211 to Houghten et al. (1986).

[089] The present invention encompasses antibodies that bind polypeptides comprising, or alternatively consisting of, an epitope of the polypeptide having an amino acid sequence of SEQ ID NO:3228, or an epitope of the polypeptide sequence encoded by a polynucleotide sequence contained in ATCC deposit No. 97768, or encoded by a polynucleotide that hybridizes to cDNA sequence contained in ATCC deposit No. 97768 (e.g., under hybridization conditions described herein).

[090] The present invention also encompasses antibodies that bind polypeptides comprising, or alternatively consisting of, an epitope of the polypeptide having an amino acid sequence of SEQ ID NO:3229, or an epitope of the polypeptide sequence encoded by a polynucleotide sequence contained in ATCC deposit No. 203518, or encoded by a polynucleotide that hybridizes to the cDNA sequence contained in ATCC deposit No. 203518 (e.g., under hybridization conditions described herein).

[091] The term "epitopes," as used herein, refers to portions of a polypeptide having antigenic or immunogenic activity in an animal, preferably a mammal, and most preferably in a human. In a preferred embodiment, the present invention encompasses antibodies that bind a polypeptide comprising an epitope. An "immunogenic epitope," as used herein, is defined as a portion of a protein that elicits an antibody response in an animal, as determined by any method known in the art, for example, by the methods for generating antibodies described *infra*. (See, for example, Geysen et al., Proc. Natl. Acad. Sci. USA 81:3998-4002 (1983)). The term "antigenic epitope," as used herein, is defined as a portion of a protein to which an antibody can immunospecifically bind its antigen as determined by any method well known in the art, for example, by the immunoassays described herein. Immunospecific binding excludes non-specific binding but does not necessarily exclude cross-reactivity with other antigens. Antigenic epitopes need not

necessarily be immunogenic.

[092] BLyS polypeptide fragments which function as epitopes may be produced by any conventional means. (See, e.g., Houghten, Proc. Natl. Acad. Sci. USA 82:5131-5135 (1985), further described in U.S. Patent No. 4,631,211).

[093] In the present invention, antibodies of the present invention bind antigenic epitopes preferably containing a sequence of at least 4, at least 5, at least 6, at least 7, more preferably at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 20, at least 25, at least 30, at least 40, at least 50, and, most preferably, between about 15 to about 30 amino acids. Preferred polypeptides comprising immunogenic or antigenic epitopes that may be bound by antibodies of the present invention are at least 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, or 100 amino acid residues in length. Additional non-exclusive preferred antigenic epitopes include the antigenic epitopes disclosed herein, as well as portions thereof. Antigenic epitopes are useful, for example, to raise antibodies, including monoclonal antibodies, that specifically bind the epitope. Preferred antigenic epitopes include the antigenic epitopes disclosed herein, as well as any combination of two, three, four, five or more of these antigenic epitopes. Antigenic epitopes can be used as the target molecules in immunoassays. (See, for instance, Wilson et al., Cell 37:767-778 (1984); Sutcliffe et al., Science 219:660-666 (1983)).

[094] Similarly, immunogenic epitopes can be used, for example, to induce antibodies according to methods well known in the art. (See, for instance, Sutcliffe et al., supra; Wilson et al., supra; Chow et al., Proc. Natl. Acad. Sci. USA 82:910-914; and Bittle et al., J. Gen. Virol. 66:2347-2354 (1985). Preferred immunogenic epitopes include the immunogenic epitopes disclosed herein, as well as any combination of two, three, four, five or more of these immunogenic epitopes. The polypeptides comprising one or more immunogenic epitopes of BLyS may be presented for eliciting an antibody response together with a carrier protein, such as an albumin, to an animal system (such as rabbit or mouse), or, if the polypeptide is of sufficient length (at least about 25 amino acids), the polypeptide may be presented without a carrier. However, immunogenic epitopes comprising as few as 8 to 10 amino acids have been shown to be sufficient to raise antibodies capable of binding to, at the very least, linear epitopes in a denatured polypeptide (e.g., in Western blotting).

[095] Epitope-bearing BLyS polypeptides may be used to induce antibodies according to methods well known in the art including, but not limited to, in vivo immunization, in vitro immunization, and phage display methods. See, e.g., Sutcliffe et al., supra; Wilson et al., supra, and Bittle et al., J. Gen. Virol., 66:2347-2354 (1985). If in vivo immunization is used, animals may be immunized with free peptide; however, antipeptide antibody titer may be boosted by coupling the peptide to a macromolecular carrier. such as keyhole limpet hemocyanin (KLH) or tetanus toxoid. For instance, peptides containing cysteine residues may be coupled to a carrier using a linker such as maleimidobenzoyl-N-hydroxysuccinimide ester (MBS), while other peptides may be coupled to carriers using a more general linking agent such as glutaraldehyde. Animals such as rabbits, rats and mice are immunized with either free or carrier-coupled peptides, for instance, by intraperitoneal and/or intradermal injection of emulsions containing about 100 micrograms of peptide or carrier protein and Freund's adjuvant or any other adjuvant known for stimulating an immune response. Several booster injections may be needed, for instance, at intervals of about two weeks, to provide a useful titer of anti-peptide antibody which can be detected, for example, by ELISA assay using free peptide adsorbed to a solid surface. The titer of anti-peptide antibodies in serum from an immunized animal may be increased by selection of anti-peptide antibodies, for instance, by adsorption to the peptide on a solid support and elution of the selected antibodies according to methods well known in the art.

[096] As one of skill in the art will appreciate, and as discussed above, the antibodies of the present invention may bind polypeptides comprising an immunogenic or antigenic epitope fused to other polypeptide sequences. For example, the BLyS polypeptides may be fused with the constant domain of immunoglobulins (IgA, IgE, IgG, IgM), or portions thereof (CH1, CH2, CH3, or any combination thereof and portions thereof), or albumin (including but not limited to recombinant human albumin or fragments or variants thereof (see, e.g., U.S. Patent No. 5,876,969, issued March 2, 1999, EP Patent 0 413 622, and U.S. Patent No. 5,766,883, issued June 16, 1998, herein incorporated by reference in their entirety)), resulting in chimeric polypeptides. Such fusion proteins may facilitate purification and may increase half-life *in vivo*. This has been shown for chimeric proteins consisting of the first two domains of the human CD4-polypeptide and various domains of the constant regions of the heavy or light chains of

mammalian immunoglobulins. See, e.g., EP 394,827; Traunecker et al., Nature, 331:84-86 (1988). Enhanced delivery of an antigen across the epithelial barrier to the immune system has been demonstrated for antigens (e.g., insulin) conjugated to an FcRn binding partner such as IgG or Fc fragments (see, e.g., PCT Publications WO 96/22024 and WO 99/04813). IgG Fusion proteins that have a disulfide-linked dimeric structure due to the IgG portion disulfide bonds have also been found to be more efficient in binding and neutralizing other molecules than monomeric polypeptides or fragments thereof alone. See, e.g., Fountoulakis et al., J. Biochem., 270:3958-3964 (1995). Nucleic acids encoding the above epitopes can also be recombined with a gene of interest as an epitope tag (e.g., the hemagglutinin ("HA") tag or flag tag) to aid in detection and purification of the expressed polypeptide. For example, a system described by Janknecht et al. allows for the ready purification of non-denatured fusion proteins expressed in human cell lines (Janknecht et al., 1991, Proc. Natl. Acad. Sci. USA 88:8972-897). In this system, the gene of interest is subcloned into a vaccinia recombination plasmid such that the open reading frame of the gene is translationally fused to an amino-terminal tag consisting of six histidine residues. The tag serves as a matrix-binding domain for the fusion protein. Extracts from cells infected with the recombinant vaccinia virus are loaded onto Ni²⁺ nitriloacetic acid-agarose column and histidine-tagged proteins can be selectively eluted with imidazole-containing buffers.

[097] In another embodiment, the antibodies of the present invention bind BLyS polypeptides and/or the epitope-bearing fragments thereof that are fused with a heterologous antigen (e.g., polypeptide, carbohydrate, phospholipid, or nucleic acid). In specific embodiments, the heterologous antigen is an immunogen.

[098] In a more specific embodiment, the heterologous antigen is the gp120 protein of HIV, or a fragment thereof.

[099] In another embodiment, antibodies of the present invention bind BLyS polypeptides and/or the epitope-bearing fragments thereof that are fused with polypeptide sequences of another TNF ligand family member (or biologically active fragments or variants thereof). In a specific embodiment, the antibodies of the present invention bind BLyS polypeptides of the present invention are fused with a CD40L polypeptide sequence. In a preferred embodiment, the CD40L polypeptide sequence is soluble.

[0100] In another embodiment, antibodies of the present invention bind mutant

BLyS polypeptides that have been generated by random mutagenesis of a polynucleotide encoding the BLyS polypeptide, by error-prone PCR, random nucleotide insertion or other methods prior to recombination. In another embodiment, antibodies of the present invention bind one or more components, motifs, sections, parts, domains, fragments, etc., of BLyS recombined with one or more components, motifs, sections, parts, domains, fragments, etc. of one or more heterologous molecules. In preferred embodiments, the heterologous molecules are, for example, TNF-alpha, lymphotoxin-alpha (LT-alpha, also known as TNF-beta), LT-beta (found in complex heterotrimer LT-alpha2-beta), OPGL, FasL, CD27L, CD30L, CD40L, 4-1BBL, DcR3, OX40L, TNF-gamma (International Publication No. WO 96/14328), AIM-I (International Publication No. WO 97/33899), AIM-II (International Publication No. WO 97/34911), APRIL (J. Exp. Med. 188(6):1185-1190), endokine-alpha (International Publication No. WO 98/07880), OPG, OX40, and nerve growth factor (NGF), and soluble forms of Fas, CD30, CD27, CD40 and 4-IBB, TR2 (International Publication No. WO 96/34095), DR3 (International Publication No. WO 97/33904), DR4 (International Publication No. WO 98/32856), TR5 (International Publication No. WO 98/30693), TR6 (International Publication No. WO 98/30694), TR7 (International Publication No. WO 98/41629), TRANK, TR9 (International Publication No. WO 98/56892), TR10 (International Publication No. WO 98/54202),312C2 (International Publication No. WO 98/06842), TR12, CAD, and v-FLIP. In further embodiments, the heterologous molecules are any member of the TNF family. [0101] In another preferred embodiment, antibodies of the present invention bind BLyS polypeptides of the invention (including biologically active fragments or variants thereof), that are fused with soluble APRIL polypeptides (e.g., amino acid residues 105 through 250 of SEQ ID NO:3239), or biologically active fragments or variants thereof. [0102] To improve or alter the characteristics of BLyS polypeptides, protein engineering may be employed. Recombinant DNA technology known to those skilled in the art can be used to create novel mutant proteins or "muteins including single or multiple amino acid substitutions, deletions, additions or fusion proteins. Such modified

polypeptides can show, e.g., enhanced activity or increased stability. In addition, they

many proteins, including the extracellular domain or the mature form(s) of a secreted

may be purified in higher yields and show better solubility than the corresponding natural polypeptide, at least under certain purification and storage conditions. For instance, for

protein, it is known in the art that one or more amino acids may be deleted from the N-terminus or C-terminus without substantial loss of biological function. For instance, Ron et al., J. Biol. Chem., 268:2984-2988 (1993) reported modified KGF proteins that had heparin binding activity even if 3, 8, or 27 amino-terminal amino acid residues were missing. Accordingly, antibodies of the present invention may bind BLyS polypeptide mutants or variants generated by protein engineering.

In the present case, since the protein of the invention is a member of the [0103] TNF polypeptide family, deletions of N-terminal amino acids up to the Gly (G) residue at position 191 in SEQ ID NO:3228 may retain some biological activity such as, for example, the ability to stimulate lymphocyte (e.g., B cell) proliferation, differentiation, and/or activation, and cytotoxicity to appropriate target cells. Polypeptides having further N-terminal deletions including the Gly (G) residue would not be expected to retain biological activities because it is known that this residue in TNF-related polypeptides is in the beginning of the conserved domain required for biological activities. However, even if deletion of one or more amino acids from the N-terminus of a protein results in modification or loss of one or more biological functions of the protein, other functional activities may still be retained. Thus, the ability of the shortened protein to induce and/or bind to antibodies which recognize the complete or extracellular domain of the protein generally will be retained when less than the majority of the residues of the complete or extracellular domain of the protein are removed from the N-terminus. Whether a particular polypeptide lacking N-terminal residues of a complete protein retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art.

[0104] Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues deleted from the amino terminus of the amino acid sequence of the BLyS of SEQ ID NO:3228, up to the glycine residue at position 191 (Gly-191 residue from the amino terminus). In particular, the present invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, the amino acid sequence of residues n¹-285 of SEQ ID NO:3228, where n¹ is an integer in the range of the amino acid position of amino acid residues 2-190 of the amino acid sequence in SEQ ID NO:3228. More in particular, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group

consisting of residues 2-285, 3-285, 4-285, 5-285, 6-285, 7-285, 8-285, 9-285, 10-285, 11-285, 12-285, 13-285, 14-285, 15-285, 16-285, 17-285, 18-285, 19-285, 20-285, 21-285, 22-285, 23-285, 24-285, 25-285, 26-285, 27-285, 28-285, 29-285, 30-285, 31-285, 32-285, 33-285, 34-285, 35-285, 36-285, 37-285, 38-285, 39-285, 40-285, 41-285, 42-285, 43-285, 44-285, 45-285, 46-285, 47-285, 48-285, 49-285, 50-285, 51-285, 52-285, 53-285, 54-285, 55-285, 56-285, 57-285, 58-285, 59-285, 60-285, 61-285, 62-285, 63-285, 64-285, 65-285, 66-285, 67-285, 68-285, 69-285, 70-285, 71-285, 72-285, 73-285, 74-285, 75-285, 76-285, 77-285, 78-285, 79-285, 80-285, 81-285, 82-285, 83-285, 84-285, 85-285, 86-285, 87-285, 88-285, 89-285, 90-285, 91-285, 92-285, 93-285, 94-285, 95-285, 96-285, 97-285, 98-285, 99-285, 100-285, 101-285, 102-285, 103-285, 104-285, 105-285, 106-285, 107-285, 108-285, 109-285, 110-285, 111-285, 112-285, 113-285, 114-285, 115-285, 116-285, 117-285, 118-285, 119-285, 120-285, 121-285, 122-285, 123-285, 124-285, 125-285, 126-285, 127-285, 128-285, 129-285, 130-285, 131-285, 132-285, 133-285, 134-285, 135-285, 136-285, 137-285, 138-285, 139-285, 140-285, 141-285, 142-285, 143-285, 144-285, 145-285, 146-285, 147-285, 148-285, 149-285, 150-285, 151-285, 152-285, 153-285, 154-285, 155-285, 156-285, 157-285, 158-285, 159-285, 160-285, 161-285, 162-285, 163-285, 164-285, 165-285, 166-285, 167-285, 168-285, 169-285, 170-285, 171-285, 172-285, 173-285, 174-285, 175-285, 176-285, 177-285, 178-285, 179-285, 180-285, 181-285, 182-285, 183-285, 184-285, 185-285, 186-285, 187-285, 188-285, 189-285, and 190-285 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0105] Furthermore, since the predicted extracellular domain of the BLyS polypeptides of the invention may itself elicit biological activity, deletions of N- and C-terminal amino acid residues from the predicted extracellular region of the polypeptide (spanning positions Gln-73 to Leu-285 of SEQ ID NO:3228) may retain some biological activity such as, for example, ligand binding, stimulation of lymphocyte (e.g., B cell) proliferation, differentiation, and/or activation, and modulation of cell replication or modulation of target cell activities. However, even if deletion of one or more amino acids from the N-terminus of the predicted extracellular domain of a BLyS polypeptide results

in modification or loss of one or more biological functions of the polypeptide, other functional activities may still be retained. Thus, the ability of the shortened polypeptides to induce and/or bind to antibodies which recognize the complete or mature or extracellular domains of the polypeptides generally will be retained when less than the majority of the residues of the complete or mature or extracellular domains of the polypeptides are removed from the N-terminus. Whether a particular polypeptide lacking N-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art. Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues deleted from the amino terminus of the amino acid sequence of BLyS shown in SEQ ID NO:3228, up to the glycine residue at position number 280. In particular, the present invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, the amino acid sequence of residues n²-285 of SEQ ID NO:3228, where n² is an integer in the range of the amino acid position of amino acid residues 73-280 in SEQ ID NO:3228, and 73 is the position of the first residue from the N-terminus of the predicted extracellular domain of the BLyS polypeptide (disclosed in SEO ID NO:3228). More in particular, in certain embodiments, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues of O-73 to L-285; G-74 to L-285; D-75 to L-285; L-76 to L-285; A-77 to L-285; S-78 to L-285; L-79 to L-285; R-80 to L-285; A-81 to L-285; E-82 to L-285; L-83 to L-285; Q-84 to L-285; G-85 to L-285; H-86 to L-285; H-87 to L-285; A-88 to L-285; E-89 to L-285; K-90 to L-285; L-91 to L-285; P-92 to L-285; A-93 to L-285; G-94 to L-285; A-95 to L-285; G-96 to L-285; A-97 to L-285; P-98 to L-285; K-99 to L-285; A-100 to L-285; G-101 to L-285; L-102 to L-285; E-103 to L-285; E-104 to L-285; A-105 to L-285; P-106 to L-285; A-107 to L-285; V-108 to L-285; T-109 to L-285; A-110 to L-285; G-111 to L-285; L-112 to L-285; K-113 to L-285; I-114 to L-285; F-115 to L-285; E-116 to L-285; P-117 to L-285; P-118 to L-285; A-119 to L-285; P-120 to L-285; G-121 to L-285; E-122 to L-285; G-123 to L-285; N-124 to L-285; S-125 to L-285; S-126 to L-285; Q-127 to L-285; N-128 to L-285; S-129 to L-285; R-130 to L-285; N-131 to L-285; K-132 to L-285; R-133 to L-285; A-134 to L-285; V-135 to L-285; O-136 to L-285; G-137 to L-285; P-138 to L-285; E-139 to L-285; E-140 to L-285; T-141 to L-285; V-142 to L-285;

T-143 to L-285; Q-144 to L-285; D-145 to L-285; C-146 to L-285; L-147 to L-285; Q-148 to L-285; L-149 to L-285; I-150 to L-285; A-151 to L-285; D-152 to L-285; S-153 to L-285; E-154 to L-285; T-155 to L-285; P-156 to L-285; T-157 to L-285; I-158 to L-285; Q-159 to L-285; K-160 to L-285; G-161 to L-285; S-162 to L-285; Y-163 to L-285; T-164 to L-285; F-165 to L-285; V-166 to L-285; P-167 to L-285; W-168 to L-285; L-169 to L-285; L-170 to L-285; S-171 to L-285; F-172 to L-285; K-173 to L-285; R-174 to L-285: G-175 to L-285; S-176 to L-285; A-177 to L-285; L-178 to L-285; E-179 to L-285; E-180 to L-285; K-181 to L-285; E-182 to L-285; N-183 to L-285; K-184 to L-285; I-185 to L-285; L-186 to L-285; V-187 to L-285; K-188 to L-285; E-189 to L-285; T-190 to L-285; G-191 to L-285; Y-192 to L-285; F-193 to L-285; F-194 to L-285; I-195 to L-285; Y-196 to L-285; G-197 to L-285; Q-198 to L-285; V-199 to L-285; L-200 to L-285; Y-201 to L-285; T-202 to L-285; D-203 to L-285; K-204 to L-285; T-205 to L-285; Y-206 to L-285; A-207 to L-285; M-208 to L-285; G-209 to L-285; H-210 to L-285; L-211 to L-285; I-212 to L-285; Q-213 to L-285; R-214 to L-285; K-215 to L-285; K-216 to L-285; V-217 to L-285; H-218 to L-285; V-219 to L-285; F-220 to L-285; G-221 to L-285; D-222 to L-285; E-223 to L-285; L-224 to L-285; S-225 to L-285; L-226 to L-285; V-227 to L-285; T-228 to L-285; L-229 to L-285; F-230 to L-285; R-231 to L-285; C-232 to L-285; I-233 to L-285; O-234 to L-285; N-235 to L-285; M-236 to L-285; P-237 to L-285; E-238 to L-285; T-239 to L-285; L-240 to L-285; P-241 to L-285; N-242 to L-285; N-243 to L-285; S-244 to L-285; C-245 to L-285; Y-246 to L-285; S-247 to L-285; A-248 to L-285; G-249 to L-285; I-250 to L-285; A-251 to L-285; K-252 to L-285; L-253 to L-285; E-254 to L-285; E-255 to L-285; G-256 to L-285; D-257 to L-285; E-258 to L-285; L-259 to L-285; Q-260 to L-285; L-261 to L-285; A-262 to L-285; I-263 to L-285; P-264 to L-285; R-265 to L-285; E-266 to L-285; N-267 to L-285; A-268 to L-285; Q-269 to L-285; I-270 to L-285; S-271 to L-285; L-272 to L-285; D-273 to L-285; G-274 to L-285; D-275 to L-285; V-276 to L-285; T-277 to L-285; F-278 to L-285; F-279 to L-285; and G-280 to L-285 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0107] Highly preferred embodiments of the invention are directed to antibodies that bind polypeptides comprising, or alternatively consisting of, a polypeptide having an

amino acid sequence least 80%, 85%, 90% identical and more preferably at least 95%, 96%, 97%, 98%, 99% or 100% identical to BLyS polypeptide having the amino acid sequence at positions 134-285 of SEQ ID NO:3228.

[0108] Preferred embodiments of the invention are directed to antibodies that bind polypeptides comprising, or alternatively consisting of, a polypeptide having an amino acid sequence at least 90% identical to a BLyS polypeptide having the amino acid sequence at positions 134-285 of SEQ ID NO:3228. More preferred embodiments of the invention are directed to antibodies that bind polypeptides comprising, or alternatively consisting of, a polypeptide having an amino acid sequence at least 95% identical to a BLyS polypeptide having the amino acid sequence at positions 134-285 of SEQ ID NO:3228. More preferred embodiments of the invention are directed to antibodies that bind polypeptides comprising, or alternatively consisting of, a polypeptide having an amino acid sequence at least 96% identical to a BLyS polypeptide having the amino acid sequence at positions 134-285 of SEQ ID NO:3228.

[0109] Additionally, more preferred embodiments of the invention are directed to antibodies that bind polypeptides comprising, or alternatively consisting of, a polypeptide having an amino acid sequence at least 97% to a BLyS polypeptide having the amino acid sequence at positions 134-285 of SEQ ID NO:3228. Additionally, more preferred embodiments of the invention are directed to antibodies that bind polypeptides comprising, or alternatively consisting of, a polypeptide having an amino acid sequence at least 98% to a BLyS polypeptide having the amino acid sequence at positions 134-285 of SEQ ID NO:3228. Additionally, more preferred embodiments of the invention are directed to antibodies that bind polypeptides comprising, or alternatively consisting of, a polypeptide having an amino acid sequence at least 99% identical to BLyS polypeptide having the amino acid sequence at positions 134-285 of SEQ ID NO:3228.

[0110] In specific embodiments, antibodies of the present invention bind polypeptides comprising, or alternatively consisting of, one of the following N-terminally deleted polypeptide fragments of BLyS: amino acid residues Ala-71 through Leu-285, amino acid residues Ala-81 through Leu-285, amino acid residues Leu-112 through Leu-285, amino acid residues Ala-134 through Leu-285, amino acid residues Leu-147 through Leu-285, and amino acid residues Gly-161 through Leu-285 of SEQ ID NO:3228.

[0111] Similarly, many examples of biologically functional C-terminal deletion

polypeptides are known. For instance, Interferon gamma shows up to ten times higher activities by deleting 8-10 amino acid residues from the carboxy terminus of the protein (Döbeli et al., J. Biotechnology 7:199-216 (1988). Since the present protein is a member of the TNF polypeptide family, deletions of C-terminal amino acids up to the leucine residue at position 284 are expected to retain most if not all biological activity such as, for example, ligand binding, the ability to stimulate lymphocyte (e.g., B cell) proliferation, differentiation, and/or activation, and modulation of cell replication. Polypeptides having deletions of up to about 10 additional C-terminal residues (i.e., up to the glycine residue at position 274) also may retain some activity such as receptor binding, although such polypeptides would lack a portion of the conserved TNF domain which extends to about Leu-284 of SEQ ID NO:3228. However, even if deletion of one or more amino acids from the C-terminus of a protein results in modification or loss of one or more biological functions of the protein, other functional activities may still be retained. Thus, the ability of the shortened protein to induce and/or bind to antibodies which recognize the complete or mature protein generally will be retained when less than the majority of the residues of the complete or mature protein are removed from the C-terminus. Whether a particular polypeptide lacking C-terminal residues of a complete protein retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art.

[0112] Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues deleted from the carboxy terminus of the amino acid sequence of the BLyS polypeptide of SEQ ID NO:3228, up to the glycine residue at position 274 (Gly-274). In particular, the present invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, the amino acid sequence of residues 1-m¹ of the amino acid sequence in SEQ ID NO:3228, where m¹ is any integer in the range of the amino acid position of amino acid residues 274-284 in SEQ ID NO:3228. More in particular, the invention provides antibodies that bind BLyS polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues 1-274, 1-275, 1-276, 1-277, 1-278, 1-279, 1-280, 1-281, 1-282, 1-283 and 1-284 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or

99% identical to the amino acid sequence of BLyS polypeptides described above.

[0113] Also provided are antibodies that bind BLyS polypeptides comprising, or alternatively consisting of, BLyS polypeptides with one or more amino acids deleted from both the amino and the carboxyl termini, which may be described generally as having residues n¹-m¹ of SEQ ID NO:3228, where n¹ and m¹ are integers as defined above. Also included are antibodies that bind a polypeptide comprising, or alternatively consisting of, a portion of the complete BLyS amino acid sequence encoded by the deposited cDNA clone contained in ATCC Accession No. 97768 where this portion excludes from 1 to 190 amino acids from the amino terminus or from 1 to 11 amino acids from the C-terminus of the complete amino acid sequence (or any combination of these N-terminal and C-terminal deletions) encoded by the cDNA clone in the deposited plasmid.

[0114] Similarly, deletions of C-terminal amino acid residues of the predicted extracellular domain of BLyS up to the leucine residue at position 79 of SEQ ID NO:3228 may retain some biological activity, such as, for example, ligand binding, stimulation of lymphocyte (e.g., B cell) proliferation, differentiation, and/or activation, and modulation of cell replication or modulation of target cell activities. Polypeptides having further C-terminal deletions including Leu-79 of SEQ ID NO:3228 would not be expected to retain biological activities.

[0115] However, even if deletion of one or more amino acids from the C-terminus of a polypeptide results in modification or loss of one or more biological functions of the polypeptide, other functional activities may still be retained. Thus, the ability of the shortened polypeptide to induce and/or bind to antibodies which recognize the complete, mature or extracellular forms of the polypeptide generally will be retained when less than the majority of the residues of the complete, mature or extracellular forms of the polypeptide are removed from the C-terminus. Whether a particular polypeptide lacking C-terminal residues of the predicted extracellular domain retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art.

[0116] Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues deleted from the carboxy terminus of the amino acid sequence of the predicted extracellular domain of BLyS polypeptide shown in SEQ ID NO:3228, up to the leucine residue at position 79 of SEQ ID NO:3228. In particular,

the present invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, the amino acid sequence of residues 73-m² of the amino acid sequence in SEQ ID NO:3228, where m² is any integer in the range of the amino acid position of amino acid residues 79 to 285 in the amino acid sequence in SEQ ID NO:3228, and residue 78 is the position of the first residue at the C-terminus of the predicted extracellular domain of the BLyS polypeptide (disclosed in SEQ ID NO:3228). More in particular, in certain embodiments, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues Q-73 to Leu-285; Q-73 to L-284; Q-73 to K-283; Q-73 to L-282; Q-73 to A-281; Q-73 to G-280; Q-73 to F-279; Q-73 to F-278; Q-73 to T-277; Q-73 to V-276; Q-73 to D-275; Q-73 to G-274; Q-73 to D-273; Q-73 to L-272; Q-73 to S-271; Q-73 to I-270; Q-73 to Q-269; Q-73 to A-268; Q-73 to N-267; Q-73 to E-266; Q-73 to R-265; Q-73 to P-264; Q-73 to I-263; Q-73 to A-262; Q-73 to L-261; Q-73 to Q-260; Q-73 to L-259; Q-73 to E-258; Q-73 to D-257; Q-73 to G-256; Q-73 to E-255; Q-73 to E-254; Q-73 to L-253; Q-73 to K-252; Q-73 to A-251; Q-73 to I-250; Q-73 to G-249; Q-73 to A-248; Q-73 to S-247; Q-73 to Y-246; Q-73 to C-245; Q-73 to S-244; Q-73 to N-243; Q-73 to N-242; Q-73 to P-241; Q-73 to L-240; Q-73 to T-239; Q-73 to E-238; Q-73 to P-237; Q-73 to M-236; Q-73 to N-235; Q-73 to Q-234; Q-73 to I-233; Q-73 to C-232; Q-73 to R-231; Q-73 to F-230; Q-73 to L-229; Q-73 to T-228; Q-73 to V-227; Q-73 to L-226; Q-73 to S-225; Q-73 to L-224; Q-73 to E-223; Q-73 to D-222; Q-73 to G-221; Q-73 to F-220; Q-73 to V-219; Q-73 to H-218; Q-73 to V-217; Q-73 to K-216; Q-73 to K-215; Q-73 to R-214; Q-73 to Q-213; Q-73 to I-212; Q-73 to L-211; Q-73 to H-210; Q-73 to G-209; Q-73 to M-208; Q-73 to A-207; Q-73 to Y-206; Q-73 to T-205; Q-73 to K-204; Q-73 to D-203; Q-73 to T-202; Q-73 to Y-201; Q-73 to L-200; Q-73 to V-199; Q-73 to Q-198; Q-73 to G-197; Q-73 to Y-196; Q-73 to I-195; Q-73 to F-194; Q-73 to F-193; Q-73 to Y-192; Q-73 to G-191; Q-73 to T-190; Q-73 to E-189; Q-73 to K-188; Q-73 to V-187; Q-73 to L-186; Q-73 to I-185; Q-73 to K-184; Q-73 to N-183; Q-73 to E-182; Q-73 to K-181; Q-73 to E-180; Q-73 to E-179; Q-73 to L-178; Q-73 to A-177; Q-73 to S-176; Q-73 to G-175; Q-73 to R-174; Q-73 to K-173; Q-73 to F-172; Q-73 to S-171; Q-73 to L-170; Q-73 to L-169; Q-73 to W-168; Q-73 to P-167; Q-73 to V-166; Q-73 to F-165; Q-73 to T-164; Q-73 to Y-163; Q-73 to S-162; Q-73 to G-161; Q-73 to K-160; Q-73 to Q-159; Q-73 to I-158; Q-73 to T-157; Q-73 to P-156;

Q-73 to T-155; Q-73 to E-154; Q-73 to S-153; Q-73 to D-152; Q-73 to A-151; Q-73 to I-150; Q-73 to L-149; Q-73 to Q-148; Q-73 to L-147; Q-73 to C-146; Q-73 to D-145; Q-73 to Q-144; Q-73 to T-143; Q-73 to V-142; Q-73 to T-141; Q-73 to E-140; Q-73 to E-139; Q-73 to P-138; Q-73 to G-137; Q-73 to Q-136; Q-73 to V-135; Q-73 to A-134; Q-73 to R-133; Q-73 to K-132; Q-73 to N-131; Q-73 to R-130; Q-73 to S-129; Q-73 to N-128; Q-73 to Q-127; Q-73 to S-126; Q-73 to S-125; Q-73 to N-124; Q-73 to G-123; Q-73 to E-122; Q-73 to G-121; Q-73 to P-120; Q-73 to A-119; Q-73 to P-118; Q-73 to P-117; Q-73 to E-116; Q-73 to F-115; Q-73 to I-114; Q-73 to K-113; Q-73 to L-112; Q-73 to G-111; Q-73 to A-110; Q-73 to T-109; Q-73 to V-108; Q-73 to A-107; Q-73 to P-106; Q-73 to A-105; Q-73 to E-104; Q-73 to E-103; Q-73 to L-102; Q-73 to G-101; Q-73 to A-100; Q-73 to K-99; Q-73 to P-98; Q-73 to A-97; Q-73 to G-96; Q-73 to A-95; Q-73 to G-94; O-73 to A-93; O-73 to P-92; O-73 to L-91; O-73 to K-90; O-73 to E-89; O-73 to A-88; Q-73 to H-87; Q-73 to H-86; Q-73 to G-85; Q-73 to Q-84; Q-73 to L-83; Q-73 to E-82; Q-73 to A-81; Q-73 to R-80; and Q-73 to L-79 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0117] The invention also provides antibodies that bind polypeptides having one or more amino acids deleted from both the amino and the carboxyl termini of the predicted extracellular domain of BLyS, which may be described generally as having residues n^2 -m² of SEQ ID NO:3228 where n^2 and m^2 are integers as defined above.

[0118] In another embodiment, antibodies of the present invention bind polypeptides consisting of a portion of the extracellular domain of the BLyS amino acid sequence encoded by the cDNA plasmid contained in the deposit having ATCC accession no. 97768, where this portion excludes from 1 to about 206 amino acids from the amino terminus of the extracellular domain of the amino acid sequence encoded by the cDNA plasmid contained in the deposit having ATCC accession no. 97768, or from 1 to about 206 amino acids from the carboxy terminus of the extracellular domain of the amino acid sequence encoded by the cDNA plasmid contained in the deposit having ATCC accession no. 97768, or any combination of the above amino terminal and carboxy terminal deletions, of the entire extracellular domain of the amino acid sequence encoded by the

cDNA plasmid contained in the deposit having ATCC accession no. 97768.

N-terminus of a polypeptide results in modification or loss of one or more functional activities (e.g., biological activity) of the polypeptide, other functions or biological activities may still be retained. Thus, the ability of a shortened BLyS mutein to induce and/or bind to antibodies which recognize the full-length or mature forms or the extracellular domain of the polypeptide generally will be retained when less than the majority of the residues of the full-length or mature or extracellular domain of the polypeptide are removed from the N-terminus. Whether a particular polypeptide lacking N-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art. It is not unlikely that a BLyS mutein with a large number of deleted N-terminal amino acid residues may retain some functional (e.g., biological or immunogenic) activities. In fact, peptides composed of as few as six BLyS amino acid residues may often evoke an immune response.

[0120] Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues deleted from the amino terminus of the predicted full-length amino acid sequence of the BLyS shown in SEQ ID NO:3228, up to the glycine residue at position number 280 of the sequence shown SEQ ID NO:3228 and polynucleotides encoding such polypeptides. In particular, the present invention provides antibodies that bind polypeptides comprising the amino acid sequence of residues n³-285 of the sequence shown in SEQ ID NO:3228, where n³ is an integer in the range of the amino acid position of amino acid residues 1 to 280 of the amino acid sequence in SEQ ID NO:3228.

[0121] More in particular, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues of D-2 to L-285; D-3 to L-285; S-4 to L-285; T-5 to L-285; E-6 to L-285; R-7 to L-285; E-8 to L-285; Q-9 to L-285; S-10 to L-285; R-11 to L-285; L-12 to L-285; T-13 to L-285; S-14 to L-285; C-15 to L-285; L-16 to L-285; K-17 to L-285; K-18 to L-285; R-19 to L-285; E-20 to L-285; E-21 to L-285; M-22 to L-285; K-23 to L-285; L-24 to L-285; K-25 to L-285; E-26 to L-285; C-27 to L-285; V-28 to L-285; S-29 to L-285; I-30 to L-285; L-31 to L-285; P-32 to L-285; R-33 to L-285; K-34 to L-285; E-35

to L-285; S-36 to L-285; P-37 to L-285; S-38 to L-285; V-39 to L-285; R-40 to L-285; S-41 to L-285; S-42 to L-285; K-43 to L-285; D-44 to L-285; G-45 to L-285; K-46 to L-285; L-47 to L-285; L-48 to L-285; A-49 to L-285; A-50 to L-285; T-51 to L-285; L-52 to L-285; L-53 to L-285; L-54 to L-285; A-55 to L-285; L-56 to L-285; L-57 to L-285; S-58 to L-285; C-59 to L-285; C-60 to L-285; L-61 to L-285; T-62 to L-285; V-63 to L-285; V-64 to L-285; S-65 to L-285; F-66 to L-285; Y-67 to L-285; Q-68 to L-285; V-69 to L-285; A-70 to L-285; A-71 to L-285; L-72 to L-285; Q-73 to L-285; G-74 to L-285; D-75 to L-285; L-76 to L-285; A-77 to L-285; S-78 to L-285; L-79 to L-285; R-80 to L-285; A-81 to L-285; E-82 to L-285; L-83 to L-285; Q-84 to L-285; G-85 to L-285; H-86 to L-285; H-87 to L-285; A-88 to L-285; E-89 to L-285; K-90 to L-285; L-91 to L-285; P-92 to L-285; A-93 to L-285; G-94 to L-285; A-95 to L-285; G-96 to L-285; A-97 to L-285; P-98 to L-285; K-99 to L-285; A-100 to L-285; G-101 to L-285; L-102 to L-285; E-103 to L-285; E-104 to L-285; A-105 to L-285; P-106 to L-285; A-107 to L-285; V-108 to L-285; T-109 to L-285; A-110 to L-285; G-111 to L-285; L-112 to L-285; K-113 to L-285; I-114 to L-285; F-115 to L-285; E-116 to L-285; P-117 to L-285; P-118 to L-285; A-119 to L-285; P-120 to L-285; G-121 to L-285; E-122 to L-285; G-123 to L-285; N-124 to L-285; S-125 to L-285; S-126 to L-285; O-127 to L-285; N-128 to L-285; S-129 to L-285; R-130 to L-285; N-131 to L-285; K-132 to L-285; R-133 to L-285; A-134 to L-285; V-135 to L-285; Q-136 to L-285; G-137 to L-285; P-138 to L-285; E-139 to L-285; E-140 to L-285; T-141 to L-285; V-142 to L-285; T-143 to L-285; Q-144 to L-285; D-145 to L-285; C-146 to L-285; L-147 to L-285; Q-148 to L-285; L-149 to L-285; I-150 to L-285; A-151 to L-285; D-152 to L-285; S-153 to L-285; E-154 to L-285; T-155 to L-285; P-156 to L-285; T-157 to L-285; I-158 to L-285; Q-159 to L-285; K-160 to L-285; G-161 to L-285; S-162 to L-285; Y-163 to L-285; T-164 to L-285; F-165 to L-285; V-166 to L-285; P-167 to L-285; W-168 to L-285; L-169 to L-285; L-170 to L-285; S-171 to L-285; F-172 to L-285; K-173 to L-285; R-174 to L-285; G-175 to L-285; S-176 to L-285; A-177 to L-285; L-178 to L-285; E-179 to L-285; E-180 to L-285; K-181 to L-285; E-182 to L-285; N-183 to L-285; K-184 to L-285; I-185 to L-285; L-186 to L-285; V-187 to L-285; K-188 to L-285; E-189 to L-285; T-190 to L-285; G-191 to L-285; Y-192 to L-285; F-193 to L-285; F-194 to L-285; I-195 to L-285; Y-196 to L-285; G-197 to L-285; Q-198 to L-285; V-199 to L-285; L-200 to L-285; Y-201 to L-285; T-202 to L-285; D-203 to L-285; K-204 to L-285; T-205 to L-285; Y-206 to L-285; A-207 to

L-285; M-208 to L-285; G-209 to L-285; H-210 to L-285; L-211 to L-285; I-212 to L-285; O-213 to L-285; R-214 to L-285; K-215 to L-285; K-216 to L-285; V-217 to L-285; H-218 to L-285; V-219 to L-285; F-220 to L-285; G-221 to L-285; D-222 to L-285; E-223 to L-285; L-224 to L-285; S-225 to L-285; L-226 to L-285; V-227 to L-285; T-228 to L-285; L-229 to L-285; F-230 to L-285; R-231 to L-285; C-232 to L-285; I-233 to L-285; O-234 to L-285; N-235 to L-285; M-236 to L-285; P-237 to L-285; E-238 to L-285; T-239 to L-285; L-240 to L-285; P-241 to L-285; N-242 to L-285; N-243 to L-285; S-244 to L-285; C-245 to L-285; Y-246 to L-285; S-247 to L-285; A-248 to L-285; G-249 to L-285; I-250 to L-285; A-251 to L-285; K-252 to L-285; L-253 to L-285; E-254 to L-285; E-255 to L-285; G-256 to L-285; D-257 to L-285; E-258 to L-285; L-259 to L-285; Q-260 to L-285; L-261 to L-285; A-262 to L-285; I-263 to L-285; P-264 to L-285; R-265 to L-285; E-266 to L-285; N-267 to L-285; A-268 to L-285; Q-269 to L-285; I-270 to L-285; S-271 to L-285; L-272 to L-285; D-273 to L-285; G-274 to L-285; D-275 to L-285; V-276 to L-285; T-277 to L-285; F-278 to L-285; F-279 to L-285; and G-280 to L-285 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

Also as mentioned above, even if deletion of one or more amino acids from the C-terminus of a protein results in modification or loss of one or more functional activities (e.g., biological activity) of the protein, other functional activities may still be retained. Thus, the ability of a shortened BLyS mutein to induce and/or bind to antibodies which recognize the complete or mature form or the extracellular domain of the polypeptide generally will be retained when less than the majority of the residues of the complete or mature form or the extracellular domain of the polypeptide are removed from the C-terminus. Whether a particular polypeptide lacking C-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art. It is not unlikely that a BLyS mutein with a large number of deleted C-terminal amino acid residues may retain some functional (e.g., biological or immunogenic) activities. In fact, peptides composed of as few as six BLyS amino acid residues may often evoke an immune response.

[0123] Accordingly, the present invention further provides in another embodiment,

antibodies that bind polypeptides having one or more residues deleted from the carboxy terminus of the amino acid sequence of the BLyS shown in SEQ ID NO:3228, up to the glutamic acid residue at position number 6, and polynucleotides encoding such polypeptides. In particular, the present invention provides antibodies that bind polypeptides comprising the amino acid sequence of residues 1-m³ of SEQ ID NO:3228, where m³ is an integer in the range of the amino acid position of amino acid residues 6-284 of the amino acid sequence in SEQ ID NO:3228.

More in particular, the invention provides antibodies that bind polypeptides [0124] comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues M-1 to L-284; M-1 to K-283; M-1 to L-282; M-1 to A-281; M-1 to G-280; M-1 to F-279; M-1 to F-278; M-1 to T-277; M-1 to V-276; M-1 to D-275; M-1 to G-274; M-1 to D-273; M-1 to L-272; M-1 to S-271; M-1 to I-270; M-1 to Q-269; M-1 to A-268; M-1 to N-267; M-1 to E-266; M-1 to R-265; M-1 to P-264; M-1 to I-263; M-1 to A-262; M-1 to L-261; M-1 to Q-260; M-1 to L-259; M-1 to E-258; M-1 to D-257; M-1 to G-256; M-1 to E-255; M-1 to E-254; M-1 to L-253; M-1 to K-252; M-1 to A-251; M-1 to I-250; M-1 to G-249; M-1 to A-248; M-1 to S-247; M-1 to Y-246; M-1 to C-245; M-1 to S-244; M-1 to N-243; M-1 to N-242; M-1 to P-241; M-1 to L-240; M-1 to T-239; M-1 to E-238; M-1 to P-237; M-1 to M-236; M-1 to N-235; M-1 to Q-234; M-1 to I-233; M-1 to C-232; M-1 to R-231; M-1 to F-230; M-1 to L-229; M-1 to T-228; M-1 to V-227; M-1 to L-226; M-1 to S-225; M-1 to L-224; M-1 to E-223; M-1 to D-222; M-1 to G-221; M-1 to F-220; M-1 to V-219; M-1 to H-218; M-1 to V-217; M-1 to K-216; M-1 to K-215; M-1 to R-214; M-1 to Q-213; M-1 to I-212; M-1 to L-211; M-1 to H-210; M-1 to G-209; M-1 to M-208; M-1 to A-207; M-1 to Y-206; M-1 to T-205; M-1 to K-204; M-1 to D-203; M-1 to T-202; M-1 to Y-201; M-1 to L-200; M-1 to V-199; M-1 to Q-198; M-1 to G-197; M-1 to Y-196; M-1 to I-195; M-1 to F-194; M-1 to F-193; M-1 to Y-192; M-1 to G-191; M-1 to T-190; M-1 to E-189; M-1 to K-188; M-1 to V-187; M-1 to L-186; M-1 to I-185; M-1 to K-184; M-1 to N-183; M-1 to E-182; M-1 to K-181; M-1 to E-180; M-1 to E-179; M-1 to L-178; M-1 to A-177; M-1 to S-176; M-1 to G-175; M-1 to R-174; M-1 to K-173; M-1 to F-172; M-1 to S-171; M-1 to L-170; M-1 to L-169; M-1 to W-168; M-1 to P-167; M-1 to V-166; M-1 to F-165; M-1 to T-164; M-1 to Y-163; M-1 to S-162; M-1 to G-161; M-1 to K-160; M-1 to Q-159; M-1 to I-158; M-1 to T-157; M-1 to P-156; M-1 to T-155; M-1 to E-154; M-1 to S-153; M-1 to D-152; M-1 to A-151; M-1 to I-150; M-1 to L-149; M-1 to

Q-148; M-1 to L-147; M-1 to C-146; M-1 to D-145; M-1 to Q-144; M-1 to T-143; M-1 to V-142; M-1 to T-141; M-1 to E-140; M-1 to E-139; M-1 to P-138; M-1 to G-137; M-1 to O-136; M-1 to V-135; M-1 to A-134; M-1 to R-133; M-1 to K-132; M-1 to N-131; M-1 to R-130; M-1 to S-129; M-1 to N-128; M-1 to Q-127; M-1 to S-126; M-1 to S-125; M-1 to N-124; M-1 to G-123; M-1 to E-122; M-1 to G-121; M-1 to P-120; M-1 to A-119; M-1 to P-118; M-1 to P-117; M-1 to E-116; M-1 to F-115; M-1 to I-114; M-1 to K-113; M-1 to L-112; M-1 to G-111; M-1 to A-110; M-1 to T-109; M-1 to V-108; M-1 to A-107; M-1 to P-106; M-1 to A-105; M-1 to E-104; M-1 to E-103; M-1 to L-102; M-1 to G-101; M-1 to A-100; M-1 to K-99; M-1 to P-98; M-1 to A-97; M-1 to G-96; M-1 to A-95; M-1 to G-94; M-1 to A-93; M-1 to P-92; M-1 to L-91; M-1 to K-90; M-1 to E-89; M-1 to A-88; M-1 to H-87; M-1 to H-86; M-1 to G-85; M-1 to Q-84; M-1 to L-83; M-1 to E-82; M-1 to A-81; M-1 to R-80; M-1 to L-79; M-1 to S-78; M-1 to A-77; M-1 to L-76; M-1 to D-75; M-1 to G-74: M-1 to O-73: M-1 to L-72; M-1 to A-71; M-1 to A-70; M-1 to V-69; M-1 to Q-68; M-1 to Y-67; M-1 to F-66; M-1 to S-65; M-1 to V-64; M-1 to V-63; M-1 to T-62; M-1 to L-61: M-1 to C-60: M-1 to C-59: M-1 to S-58; M-1 to L-57; M-1 to L-56; M-1 to A-55; M-1 to L-54; M-1 to L-53; M-1 to L-52; M-1 to T-51; M-1 to A-50; M-1 to A-49; M-1 to L-48; M-1 to L-47; M-1 to K-46; M-1 to G-45; M-1 to D-44; M-1 to K-43; M-1 to S-42; M-1 to S-41; M-1 to R-40; M-1 to V-39; M-1 to S-38; M-1 to P-37; M-1 to S-36; M-1 to E-35; M-1 to K-34; M-1 to R-33; M-1 to P-32; M-1 to L-31; M-1 to I-30; M-1 to S-29; M-1 to V-28; M-1 to C-27; M-1 to E-26; M-1 to K-25; M-1 to L-24; M-1 to K-23; M-1 to M-22; M-1 to E-21; M-1 to E-20; M-1 to R-19; M-1 to K-18; M-1 to K-17; M-1 to L-16; M-1 to C-15; M-1 to S-14; M-1 to T-13; M-1 to L-12; M-1 to R-11; M-1 to S-10; M-1 to Q-9; M-1 to E-8; M-1 to R-7; and M-1 to E-6 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0125] The invention also provides antibodies that bind polypeptides having one or more amino acids deleted from both the amino and the carboxyl termini of a BLyS polypeptide, which may be described generally as having residues n³-m³ of SEQ ID NO:3228, where n³ and m³ are integers as defined above.

[0126] Furthermore, since the predicted extracellular domain of the BLyS

polypeptide of SEQ ID NO:3229 may itself elicit functional activity (e.g., biological activity), deletions of N- and C-terminal amino acid residues from the predicted extracellular region of the polypeptide at positions Gln-73 to Leu-266 of SEQ ID NO:3229 may retain some functional activity, such as, for example, ligand binding, to stimulation of lymphocyte (e.g., B cell) proliferation, differentiation, and/or activation, modulation of cell replication, modulation of target cell activities and/or immunogenicity. However, even if deletion of one or more amino acids from the N-terminus of the predicted extracellular domain of a BLyS polypeptide results in modification or loss of one or more functional activities of the polypeptide, other functional activities may still be retained. Thus, the ability of the shortened polypeptides to induce and/or bind to antibodies which recognize the complete or mature or extracellular domains of the polypeptides generally will be retained when less than the majority of the residues of the complete or mature or extracellular domains of the polypeptides are removed from the N-terminus. Whether a particular polypeptide lacking N-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art.

[0127] Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues deleted from the amino terminus of the amino acid sequence of BLyS shown in SEQ ID NO:3229, up to the glycine residue at position number 261. In particular, the present invention provides antibodies that bind polypeptides comprising the amino acid sequence of residues n⁴-266 of SEQ ID NO:3229, where n⁴ is an integer in the range of the amino acid position of amino acid residues 73-261 of the amino acid sequence in SEQ ID NO:3229, and 261 is the position of the first residue from the N-terminus of the predicted extracellular domain BLyS polypeptide (shown in SEQ ID NO:3229).

[0128] More in particular, in certain embodiments, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues of Q-73 to L-266; G-74 to L-266; D-75 to L-266; L-76 to L-266; A-77 to L-266; S-78 to L-266; L-79 to L-266; R-80 to L-266; A-81 to L-266; E-82 to L-266; L-83 to L-266; Q-84 to L-266; G-85 to L-266; H-86 to L-266; H-87 to L-266; A-88 to L-266; E-89 to L-266; K-90 to L-266; L-91 to L-266; P-92 to L-266; A-93 to L-266; G-94 to L-266; A-95 to L-266; G-96 to L-266; A-97 to

L-266; P-98 to L-266; K-99 to L-266; A-100 to L-266; G-101 to L-266; L-102 to L-266; E-103 to L-266; E-104 to L-266; A-105 to L-266; P-106 to L-266; A-107 to L-266; V-108 to L-266; T-109 to L-266; A-110 to L-266; G-111 to L-266; L-112 to L-266; K-113 to L-266; I-114 to L-266; F-115 to L-266; E-116 to L-266; P-117 to L-266; P-118 to L-266; A-119 to L-266; P-120 to L-266; G-121 to L-266; E-122 to L-266; G-123 to L-266; N-124 to L-266; S-125 to L-266; S-126 to L-266; Q-127 to L-266; N-128 to L-266; S-129 to L-266; R-130 to L-266; N-131 to L-266; K-132 to L-266; R-133 to L-266; A-134 to L-266; V-135 to L-266; Q-136 to L-266; G-137 to L-266; P-138 to L-266; E-139 to L-266; E-140 to L-266; T-141 to L-266; G-142 to L-266; S-143 to L-266; Y-144 to L-266; T-145 to L-266; F-146 to L-266; V-147 to L-266; P-148 to L-266; W-149 to L-266; L-150 to L-266; L-151 to L-266; S-152 to L-266; F-153 to L-266; K-154 to L-266; R-155 to L-266; G-156 to L-266; S-157 to L-266; A-158 to L-266; L-159 to L-266; E-160 to L-266; E-161 to L-266; K-162 to L-266; E-163 to L-266; N-164 to L-266; K-165 to L-266; I-166 to L-266; L-167 to L-266; V-168 to L-266; K-169 to L-266; E-170 to L-266; T-171 to L-266; G-172 to L-266; Y-173 to L-266; F-174 to L-266; F-175 to L-266; I-176 to L-266; Y-177 to L-266; G-178 to L-266; Q-179 to L-266; V-180 to L-266; L-181 to L-266; Y-182 to L-266; T-183 to L-266; D-184 to L-266; K-185 to L-266; T-186 to L-266; Y-187 to L-266; A-188 to L-266; M-189 to L-266; G-190 to L-266; H-191 to L-266; L-192 to L-266; I-193 to L-266; Q-194 to L-266; R-195 to L-266; K-196 to L-266; K-197 to L-266; V-198 to L-266; H-199 to L-266; V-200 to L-266; F-201 to L-266; G-202 to L-266; D-203 to L-266; E-204 to L-266; L-205 to L-266; S-206 to L-266; L-207 to L-266; V-208 to L-266; T-209 to L-266; L-210 to L-266; F-211 to L-266; R-212 to L-266; C-213 to L-266; I-214 to L-266; Q-215 to L-266; N-216 to L-266; M-217 to L-266; P-218 to L-266; E-219 to L-266; T-220 to L-266; L-221 to L-266; P-222 to L-266; N-223 to L-266; N-224 to L-266; S-225 to L-266; C-226 to L-266; Y-227 to L-266; S-228 to L-266; A-229 to L-266; G-230 to L-266; I-231 to L-266; A-232 to L-266; K-233 to L-266; L-234 to L-266; E-235 to L-266; E-236 to L-266; G-237 to L-266; D-238 to L-266; E-239 to L-266; L-240 to L-266; Q-241 to L-266; L-242 to L-266; A-243 to L-266; I-244 to L-266; P-245 to L-266; R-246 to L-266; E-247 to L-266; N-248 to L-266; A-249 to L-266; Q-250 to L-266; I-251 to L-266; S-252 to L-266; L-253 to L-266; D-254 to L-266; G-255 to L-266; D-256 to L-266; V-257 to L-266; T-258 to L-266; F-259 to L-266; F-260 to L-266; and G-261 to L-266 of SEQ ID NO:3229. The present invention is also directed to antibodies

that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0129] Similarly, deletions of C-terminal amino acid residues of the predicted extracellular domain of BLyS up to the leucine residue at position 79 of SEQ ID NO:3229 may retain some functional activity, such as, for example, ligand binding, the ability to stimulate lymphocyte (e.g., B cell) proliferation, differentiation, and/or activation, modulation of cell replication, modulation of target cell activities and/or immunogenicity. Polypeptides having further C-terminal deletions including Leu-79 of SEQ ID NO:3229 would not be expected to retain biological activities.

[0130] However, even if deletion of one or more amino acids from the C-terminus of a polypeptide results in modification or loss of one or more functional activities (e.g., biological activity) of the polypeptide, other functional activities may still be retained. Thus, the ability of the shortened polypeptide to induce and/or bind to antibodies which recognize the complete, mature or extracellular forms of the polypeptide generally will be retained when less than the majority of the residues of the complete, mature or extracellular forms of the polypeptide are removed from the C-terminus. Whether a particular polypeptide lacking C-terminal residues of the predicted extracellular domain retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art.

[0131] Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues from the carboxy terminus of the amino acid sequence of the predicted extracellular domain of BLyS shown in SEQ ID NO:3229, up to the leucine residue at position 79 of SEQ ID NO:3229. In particular, the present invention provides antibodies that bind polypeptides having the amino acid sequence of residues 73-m⁴ of the amino acid sequence in SEQ ID NO:3229, where m⁴ is any integer in the range of the amino acid position of amino acid residues 79-265 of the amino acid sequence in SEQ ID NO:3229.

[0132] More in particular, in certain embodiments, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues Q-73 to L-265; Q-73 to K-264; Q-73 to L-263; Q-73 to A-262; Q-73 to G-261; Q-73 to F-260; Q-73 to F-259; Q-73 to

T-258; Q-73 to V-257; Q-73 to D-256; Q-73 to G-255; Q-73 to D-254; Q-73 to L-253; Q-73 to S-252; Q-73 to I-251; Q-73 to Q-250; Q-73 to A-249; Q-73 to N-248; Q-73 to E-247; Q-73 to R-246; Q-73 to P-245; Q-73 to I-244; Q-73 to A-243; Q-73 to L-242; Q-73 to Q-241; Q-73 to L-240; Q-73 to E-239; Q-73 to D-238; Q-73 to G-237; Q-73 to E-236; Q-73 to E-235; Q-73 to L-234; Q-73 to K-233; Q-73 to A-232; Q-73 to I-231; Q-73 to G-230; Q-73 to A-229; Q-73 to S-228; Q-73 to Y-227; Q-73 to C-226; Q-73 to S-225; Q-73 to N-224; Q-73 to N-223; Q-73 to P-222; Q-73 to L-221; Q-73 to T-220; Q-73 to E-219; Q-73 to P-218; Q-73 to M-217; Q-73 to N-216; Q-73 to Q-215; Q-73 to I-214; Q-73 to C-213; Q-73 to R-212; Q-73 to F-211; Q-73 to L-210; Q-73 to T-209; Q-73 to V-208; O-73 to L-207; Q-73 to S-206; Q-73 to L-205; Q-73 to E-204; Q-73 to D-203; Q-73 to G-202; Q-73 to F-201; Q-73 to V-200; Q-73 to H-199; Q-73 to V-198; Q-73 to K-197; Q-73 to K-196; Q-73 to R-195; Q-73 to Q-194; Q-73 to I-193; Q-73 to L-192; Q-73 to H-191; Q-73 to G-190; Q-73 to Q-7389; Q-73 to A-188; Q-73 to Y-187; Q-73 to T-186; Q-73 to K-185; Q-73 to D-184; Q-73 to T-183; Q-73 to Y-182; Q-73 to L-181; Q-73 to V-180; Q-73 to Q-179; Q-73 to G-178; Q-73 to Y-177; Q-73 to I-176; Q-73 to F-175; Q-73 to F-174; Q-73 to Y-173; Q-73 to G-172; Q-73 to T-171; Q-73 to E-170; Q-73 to K-169; Q-73 to V-168; Q-73 to L-167; Q-73 to I-166; Q-73 to K-165; Q-73 to N-164; Q-73 to E-163; Q-73 to K-162; Q-73 to E-161; Q-73 to E-160; Q-73 to L-159; Q-73 to A-158; Q-73 to S-157; Q-73 to G-156; Q-73 to R-155; Q-73 to K-154; Q-73 to F-153; Q-73 to S-152; Q-73 to L-151; Q-73 to L-150; Q-73 to W-149; Q-73 to P-148; Q-73 to V-147; Q-73 to F-146; Q-73 to T-145; Q-73 to Y-144; Q-73 to S-143; Q-73 to G-142; Q-73 to T-141; Q-73 to E-140; Q-73 to E-139; Q-73 to P-138; Q-73 to G-137; Q-73 to Q-136; Q-73 to V-135; Q-73 to A-134; Q-73 to R-133; Q-73 to K-132; Q-73 to N-131; Q-73 to R-130; Q-73 to S-129; Q-73 to N-128; Q-73 to Q-127; Q-73 to S-126; Q-73 to S-125; Q-73 to N-124; Q-73 to G-123; Q-73 to E-122; Q-73 to G-121; Q-73 to P-120; Q-73 to A-119; Q-73 to P-118; Q-73 to P-117; Q-73 to E-116; Q-73 to F-115; Q-73 to I-114; Q-73 to K-113; Q-73 to L-112; Q-73 to G-111; Q-73 to A-110; Q-73 to T-109; Q-73 to V-108; Q-73 to A-107; Q-73 to P-106; Q-73 to A-105; Q-73 to E-104; Q-73 to E-103; Q-73 to L-102; Q-73 to G-101; Q-73 to A-100; Q-73 to K-99; Q-73 to P-98; Q-73 to A-97; Q-73 to G-96; Q-73 to A-95; Q-73 to G-94; Q-73 to A-93; Q-73 to P-92; Q-73 to L-91; Q-73 to K-90; Q-73 to E-89; Q-73 to A-88; Q-73 to H-87; Q-73 to H-86; Q-73 to G-85; Q-73 to Q-84; Q-73 to L-83; Q-73 to E-82; Q-73 to A-81; Q-73 to

R-80; Q-73 to L-79; and Q-73 to S-78 of SEQ ID NO:3229. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0133] The invention also provides polypeptides having one or more amino acids deleted from both the amino and the carboxyl termini of the predicted extracellular domain of BLyS, which may be described generally as having residues n⁴-m⁴ of SEQ ID NO:3229 where n⁴ and m⁴ are integers as defined above.

[0134] In another embodiment, antibodies of the present invention bind polypeptides consisting of a portion of the extracellular domain of the BLyS amino acid sequence encoded by the cDNA clone contained in the deposit having ATCC Accession No. 203518, where this portion excludes from 1 to about 260 amino acids from the amino terminus of the extracellular domain of the amino acid sequence encoded by cDNA clone contained in the deposit having ATCC Accession No. 203518, or from 1 to about 187 amino acids from the carboxy terminus of the extracellular domain of the amino acid sequence encoded by cDNA clone contained in the deposit having ATCC Accession No. 203518, or any combination of the above amino terminal and carboxy terminal deletions, of the entire extracellular domain of the amino acid sequence encoded by the cDNA clone contained in the deposit having ATCC Accession No. 203518.

[0135] As mentioned above, even if deletion of one or more amino acids from the N-terminus of a polypeptide results in modification or loss of one or more functional activities (e.g., biological activity) of the polypeptide, other functional activities may still be retained. Thus, the ability of a shortened BLyS polypeptide to induce and/or bind to antibodies which recognize the full-length or mature forms or the extracellular domain of the polypeptide generally will be retained when less than the majority of the residues of the full-length or mature or extracellular domain of the polypeptide are removed from the N-terminus. Whether a particular polypeptide lacking N-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art. It is not unlikely that a BLyS mutein with a large number of deleted N-terminal amino acid residues may retain functional (e.g., immunogenic) activities. In fact, peptides composed of as few as six

BLyS amino acid residues may often evoke an immune response.

[0136] Accordingly, the present invention further provides antibodies that bind polypeptides having one or more residues deleted from the amino terminus of the predicted full-length amino acid sequence of the BLyS polypeptide shown in SEQ ID NO:3229, up to the glycine residue at position number 261 of the sequence shown SEQ ID NO:3229 and polynucleotides encoding such polypeptides. In particular, the present invention provides antibodies that bind polypeptides comprising the amino acid sequence of residues n⁵-266 of the sequence shown in SEQ ID NO:3229, where n⁵ is an integer in the range of the amino acid position of amino acid residues 1 to 261 of the amino acid sequence in SEQ ID NO:3229.

[0137] More in particular, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues of D-2 to L-266; D-3 to L-266; S-4 to L-266; T-5 to L-266; E-6 to L-266; R-7 to L-266; E-8 to L-266; Q-9 to L-266; S-10 to L-266; R-11 to L-266; L-12 to L-266; T-13 to L-266; S-14 to L-266; C-15 to L-266; L-16 to L-266; K-17 to L-266; K-18 to:L-266; R-19 to L-266; E-20 to L-266; E-21 to L-266; M-22 to L-266; K-23 to L-266; L-24 to L-266; K-25 to L-266; E-26 to L-266; C-27 to L-266; V-28 to L-266; S-29 to L-266; I-30 to L-266; L-31 to L-266; P-32 to L-266; R-33 to L-266; K-34 to L-266; E-35 to L-266; S-36 to L-266; P-37 to L-266; S-38 to L-266; V-39 to L-266; R-40 to L-266; S-41 to L-266; S-42 to L-266; K-43 to L-266; D-44 to L-266; G-45 to L-266; K-46 to L-266; L-47 to L-266; L-48 to L-266; A-49 to L-266; A-50 to L-266; T-51 to L-266; L-52 to L-266; L-53 to L-266; L-54 to L-266; A-55 to L-266; L-56 to L-266; L-57 to L-266; S-58 to L-266; C-59 to L-266; C-60 to L-266; L-61 to L-266; T-62 to L-266; V-63 to L-266; V-64 to L-266; S-65 to L-266; F-66 to L-266; Y-67 to L-266; Q-68 to L-266; V-69 to L-266; A-70 to L-266; A-71 to L-266; L-72 to L-266; Q-73 to L-266; G-74 to L-266; D-75 to L-266; L-76 to L-266; A-77 to L-266; S-78 to L-266; L-79 to L-266; R-80 to L-266; A-81 to L-266; E-82 to L-266; L-83 to L-266; Q-84 to L-266; G-85 to L-266; H-86 to L-266; H-87 to L-266; A-88 to L-266; E-89 to L-266; K-90 to L-266; L-91 to L-266; P-92 to L-266; A-93 to L-266; G-94 to L-266; A-95 to L-266; G-96 to L-266; A-97 to L-266; P-98 to L-266; K-99 to L-266; A-100 to L-266; G-101 to L-266; L-102 to L-266; E-103 to L-266; E-104 to L-266; A-105 to L-266; P-106 to L-266; A-107 to L-266; V-108 to L-266; T-109 to L-266; A-110 to L-266; G-111 to L-266; L-112 to L-266; K-113 to

L-266; I-114 to L-266; F-115 to L-266; E-116 to L-266; P-117 to L-266; P-118 to L-266; A-119 to L-266; P-120 to L-266; G-121 to L-266; E-122 to L-266; G-123 to L-266; N-124 to L-266; S-125 to L-266; S-126 to L-266; Q-127 to L-266; N-128 to L-266; S-129 to L-266; R-130 to L-266; N-131 to L-266; K-132 to L-266; R-133 to L-266; A-134 to L-266; V-135 to L-266; Q-136 to L-266; G-137 to L-266; P-138 to L-266; E-139 to L-266; E-140 to L-266; T-141 to L-266; G-142 to L-266; S-143 to L-266; Y-144 to L-266; T-145 to L-266; F-146 to L-266; V-147 to L-266; P-148 to L-266; W-149 to L-266; L-150 to L-266; L-151 to L-266; S-152 to L-266; F-153 to L-266; K-154 to L-266; R-155 to L-266; G-156 to L-266; S-157 to L-266; A-158 to L-266; L-159 to L-266; E-160 to L-266; E-161 to L-266; K-162 to L-266; E-163 to L-266; N-164 to L-266; K-165 to L-266; I-166 to L-266; L-167 to L-266; V-168 to L-266; K-169 to L-266; E-170 to L-266; T-171 to L-266; G-172 to L-266; Y-173 to L-266; F-174 to L-266; F-175 to L-266; I-176 to L-266: Y-177 to L-266; G-178 to L-266; Q-179 to L-266; V-180 to L-266; L-181 to L-266; Y-182 to L-266; T-183 to L-266; D-184 to L-266; K-185 to L-266; T-186 to L-266; Y-187 to L-266; A-188 to L-266; M-189 to L-266; G-190 to L-266; H-191 to L-266; L-192 to L-266; I-193 to L-266; Q-194 to L-266; R-195 to L-266; K-196 to L-266; K-197 to L-266; V-198 to L-266; H-199 to L-266; V-200 to L-266; F-201 to L-266; G-202 to L-266; D-203 to L-266; E-204 to L-266; L-205 to L-266; S-206 to L-266; L-207 to L-266; V-208 to L-266; T-209 to L-266; L-210 to L-266; F-211 to L-266; R-212 to L-266; C-213 to L-266; I-214 to L-266; Q-215 to L-266; N-216 to L-266; M-217 to L-266; P-218 to L-266; E-219 to L-266; T-220 to L-266; L-221 to L-266; P-222 to L-266; N-223 to L-266; N-224 to L-266; S-225 to L-266; C-226 to L-266; Y-227 to L-266; S-228 to L-266; A-229 to L-266; G-230 to L-266; I-231 to L-266; A-232 to L-266; K-233 to L-266; L-234 to L-266; E-235 to L-266; E-236 to L-266; G-237 to L-266; D-238 to L-266; E-239 to L-266; L-240 to L-266; Q-241 to L-266; L-242 to L-266; A-243 to L-266; I-244 to L-266; P-245 to L-266; R-246 to L-266; E-247 to L-266; N-248 to L-266; A-249 to L-266; Q-250 to L-266; I-251 to L-266; S-252 to L-266; L-253 to L-266; D-254 to L-266; G-255 to L-266; D-256 to L-266; V-257 to L-266; T-258 to L-266; F-259 to L-266; F-260 to L-266; and G-261 to L-266 of SEQ ID NO:3229. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0138] Also as mentioned above, even if deletion of one or more amino acids from the C-terminus of a protein results in modification or loss of one or more functional activities (e.g., biological activities) of the protein, other functional activities may still be retained. Thus, the ability of a shortened BLyS mutein to induce and/or bind to antibodies which recognize the complete or mature form or the extracellular domain of the polypeptide generally will be retained when less than the majority of the residues of the complete or mature form or the extracellular domain of the polypeptide are removed from the C-terminus. Whether a particular polypeptide lacking C-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art. It is not unlikely that a BLyS mutein with a large number of deleted C-terminal amino acid residues may retain some functional (e.g., immunogenic) activities. In fact, peptides composed of as few as six BLyS amino acid residues may often evoke an immune response.

[0139] Accordingly, the present invention further provides in another embodiment, antibodies that bind polypeptides having one or more residues deleted from the carboxy terminus of the amino acid sequence of the BLyS shown in SEQ ID NO:3229, up to the glutamic acid residue at position number 6, and polynucleotides encoding such polypeptides. In particular, the present invention provides antibodies that bind polypeptides comprising the amino acid sequence of residues 1-m⁵ of SEQ ID NO:3229, where m⁵ is an integer in the range of the amino acid position of amino acid residues 6 to 265 in the amino acid sequence of SEQ ID NO:3229.

[0140] More in particular, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues M-1 to L-265; M-1 to K-264; M-1 to L-263; M-1 to A-262; M-1 to G-261; M-1 to F-260; M-1 to F-259; M-1 to T-258; M-1 to V-257; M-1 to D-256; M-1 to G-255; M-1 to D-254; M-1 to L-253; M-1 to S-252; M-1 to I-251; M-1 to Q-250; M-1 to A-249; M-1 to N-248; M-1 to E-247; M-1 to R-246; M-1 to P-245; M-1 to I-244; M-1 to A-243; M-1 to L-242; M-1 to Q-241; M-1 to L-240; M-1 to E-239; M-1 to D-238; M-1 to G-237; M-1 to E-236; M-1 to E-235; M-1 to L-234; M-1 to K-233; M-1 to A-232; M-1 to I-231; M-1 to G-230; M-1 to A-229; M-1 to S-228; M-1 to Y-227; M-1 to C-226; M-1 to S-225; M-1 to N-224; M-1 to N-223; M-1 to P-222; M-1 to L-221; M-1 to T-220; M-1 to E-219; M-1 to P-218; M-1 to M-217; M-1 to N-216; M-1 to Q-215; M-1 to I-214; M-1 to

C-213; M-1 to R-212; M-1 to F-211; M-1 to L-210; M-1 to T-209; M-1 to V-208; M-1 to L-207; M-1 to S-206; M-1 to L-205; M-1 to E-204; M-1 to D-203; M-1 to G-202; M-1 to F-201; M-1 to V-200; M-1 to H-199; M-1 to V-198; M-1 to K-197; M-1 to K-196; M-1 to R-195; M-1 to Q-194; M-1 to I-193; M-1 to L-192; M-1 to H-191; M-1 to G-190; M-1 to M-189; M-1 to A-188; M-1 to Y-187; M-1 to T-186; M-1 to K-185; M-1 to D-184; M-1 to T-183; M-1 to Y-182; M-1 to L-181; M-1 to V-180; M-1 to Q-179; M-1 to G-178; M-1 to Y-177; M-1 to I-176; M-1 to F-175; M-1 to F-174; M-1 to Y-173; M-1 to G-172; M-1 to T-171; M-1 to E-170; M-1 to K-169; M-1 to V-168; M-1 to L-167; M-1 to I-166; M-1 to K-165; M-1 to N-164; M-1 to E-163; M-1 to K-162; M-1 to E-161; M-1 to E-160; M-1 to L-159; M-1 to A-158; M-1 to S-157; M-1 to G-156; M-1 to R-155; M-1 to K-154; M-1 to F-153; M-1 to S-152; M-1 to L-151; M-1 to L-150; M-1 to W-149; M-1 to P-148; M-1 to V-147; M-1 to F-146; M-1 to T-145; M-1 to Y-144; M-1 to S-143; M-1 to G-142; M-1 to T-141; M-1 to E-140; M-1 to E-139; M-1 to P-138; M-1 to G-137; M-1 to Q-136; M-1 to V-135; M-1 to A-134; M-1 to R-133; M-1 to K-132; M-1 to N-131; M-1 to R-130; M-1 to S-129; M-1 to N-128; M-1 to Q-127; M-1 to S-126; M-1 to S-125; M-1 to N-124; M-1 to G-123; M-1 to E-122; M-1 to G-121; M-1 to P-120; M-1 to A-119; M-1 to P-118; M-1 to P-117; M-1 to E-116; M-1 to F-115; M-1 to I-114; M-1 to K-113; M-1 to L-112; M-1 to G-111; M-1 to A-110; M-1 to T-109; M-1 to V-108; M-1 to A-107; M-1 to P-106; M-1 to A-105; M-1 to E-104; M-1 to E-103; M-1 to L-102; M-1 to G-101; M-1 to A-100; M-1 to K-99; M-1 to P-98; M-1 to A-97; M-1 to G-96; M-1 to A-95; M-1 to G-94; M-1 to A-93; M-1 to P-92; M-1 to L-91; M-1 to K-90; M-1 to E-89; M-1 to A-88; M-1 to H-87; M-1 to H-86; M-1 to G-85; M-1 to Q-84; M-1 to L-83; M-1 to E-82; M-1 to A-81; M-1 to R-80; M-1 to L-79; M-1 to S-78; M-1 to A-77; M-1 to L-76; M-1 to D-75; M-1 to G-74; M-1 to Q-73; M-1 to L-72; M-1 to A-71; M-1 to A-70; M-1 to V-69; M-1 to O-68; M-1 to Y-67; M-1 to F-66; M-1 to S-65; M-1 to V-64; M-1 to V-63; M-1 to T-62; M-1 to L-61; M-1 to C-60; M-1 to C-59; M-1 to S-58; M-1 to L-57; M-1 to L-56; M-1 to A-55; M-1 to L-54; M-1 to L-53; M-1 to L-52; M-1 to T-51; M-1 to A-50; M-1 to A-49; M-1 to L-48; M-1 to L-47; M-1 to K-46; M-1 to G-45; M-1 to D-44; M-1 to K-43; M-1 to S-42; M-1 to S-41; M-1 to R-40; M-1 to V-39; M-1 to S-38; M-1 to P-37; M-1 to S-36; M-1 to E-35; M-1 to K-34; M-1 to R-33; M-1 to P-32; M-1 to L-31; M-1 to I-30; M-1 to S-29; M-1 to V-28; M-1 to C-27; M-1 to E-26; M-1 to K-25; M-1 to L-24; M-1 to K-23; M-1 to M-22; M-1 to E-21; M-1 to E-20; M-1 to R-19; M-1 to K-18; M-1 to K-17; M-1 to L-16; M-1 to C-15;

M-1 to S-14; M-1 to T-13; M-1 to L-12; M-1 to R-11; M-1 to S-10; M-1 to Q-9; M-1 to E-8; M-1 to R-7; and M-1 to E-6 of SEQ ID NO:3229. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0141] The invention also provides antibodies that bind polypeptides having one or more amino acids deleted from both the amino and the carboxyl termini of a BLyS polypeptide, which may be described generally as having residues n⁵-m⁵ of SEQ ID NO:3229, where n⁵ and m⁵ are integers as defined above.

[0142] In additional embodiments, the present invention provides antibodies that bind polypeptides comprising the amino acid sequence of residues 134-m⁶ of SEO ID NO:3228, where m⁶ is an integer from 140 to 285, corresponding to the position of the amino acid residue in SEQ ID NO:3228. For example, the invention provides antibodies that bind polypeptides comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues A-134 to Leu-285; A-134 to L-284; A-134 to K-283; A-134 to L-282; A-134 to A-281; A-134 to G-280; A-134 to F-279; A-134 to F-278; A-134 to T-277; A-134 to V-276; A-134 to D-275; A-134 to G-274; A-134 to D-273; A-134 to L-272; A-134 to S-271; A-134 to I-270; A-134 to Q-269; A-134 to A-268; A-134 to N-267; A-134 to E-266; A-134 to R-265; A-134 to P-264; A-134 to I-263; A-134 to A-262; A-134 to L-261; A-134 to Q-260; A-134 to L-259; A-134 to E-258; A-134 to D-257; A-134 to G-256; A-134 to E-255; A-134 to E-254; A-134 to L-253; A-134 to K-252; A-134 to A-251; A-134 to I-250; A-134 to G-249; A-134 to A-248; A-134 to S-247; A-134 to Y-246; A-134 to C-245; A-134 to S-244; A-134 to N-243; A-134 to N-242; A-134 to P-241; A-134 to L-240; A-134 to T-239; A-134 to E-238; A-134 to P-237; A-134 to M-236; A-134 to N-235; A-134 to Q-234; A-134 to I-233; A-134 to C-232; A-134 to R-231; A-134 to F-230; A-134 to L-229; A-134 to T-228; A-134 to V-227; A-134 to L-226; A-134 to S-225; A-134 to L-224; A-134 to E-223; A-134 to D-222; A-134 to G-221; A-134 to F-220; A-134 to V-219; A-134 to H-218; A-134 to V-217; A-134 to K-216; A-134 to K-215; A-134 to R-214; A-134 to Q-213; A-134 to I-212; A-134 to L-211; A-134 to H-210; A-134 to G-209; A-134 to M-208; A-134 to A-207; A-134 to Y-206; A-134 to T-205; A-134 to K-204; A-134 to

D-203; A-134 to T-202; A-134 to Y-201; A-134 to L-200; A-134 to V-199; A-134 to Q-198; A-134 to G-197; A-134 to Y-196; A-134 to I-195; A-134 to F-194; A-134 to F-193; A-134 to Y-192; A-134 to G-191; A-134 to T-190; A-134 to E-189; A-134 to K-188; A-134 to V-187; A-134 to L-186; A-134 to I-185; A-134 to K-184; A-134 to N-183; A-134 to E-182; A-134 to K-181; A-134 to E-180; A-134 to E-179; A-134 to L-178; A-134 to A-177; A-134 to S-176; A-134 to G-175; A-134 to R-174; A-134 to K-173; A-134 to F-172; A-134 to S-171; A-134 to L-170; A-134 to L-169; A-134 to W-168; A-134 to P-167; A-134 to V-166; A-134 to F-165; A-134 to T-164; A-134 to Y-163; A-134 to S-162; A-134 to G-161; A-134 to K-160; A-134 to O-159; A-134 to I-158; A-134 to T-157; A-134 to P-156; A-134 to T-155; A-134 to E-154; A-134 to S-153; A-134 to D-152; A-134 to A-151; A-134 to I-150; A-134 to L-149; A-134 to Q-148; A-134 to L-147; A-134 to C-146; A-134 to D-145; A-134 to Q-144; A-134 to T-143; A-134 to V-142; A-134 to T-141; and A-134 to E-140 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

In additional embodiments, antibodies of the present invention may bind [0143] polypeptide fragments comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues: M-1 to C-15; D-2 to L-16; D-3 to K-17; S-4 to K-18; T-5 to R-19; E-6 to E-20; R-7 to E-21; E-8 to M-22; Q-9 to K-23; S-10 to L-24; R-11 to K-25; L-12 to E-26; T-13 to C-27; S-14 to V-28; C-15 to S-29; L-16 to I-30; K-17 to L-31; K-18 to P-32; R-19 to R-33; E-20 to K-34; E-21 to E-35; M-22 to S-36; K-23 to P-37; L-24 to S-38; K-25 to V-39; E-26 to R-40; C-27 to S-41; V-28 to S-42; S-29 to K-43; I-30 to D-44; L-31 to G-45; P-32 to K-46; R-33 to L-47; K-34 to L-48; E-35 to A-49; S-36 to A-50; P-37 to T-51; S-38 to L-52; V-39 to L-53; R-40 to L-54; S-41 to A-55; S-42 to L-56; K-43 to L-57; D-44 to S-58; G-45 to C-59; K-46 to C-60; L-47 to L-61; L-48 to T-62; A-49 to V-63; A-50 to V-64; T-51 to S-65; L-52 to F-66; L-53 to Y-67; L-54 to Q-68; A-55 to V-69; L-56 to A-70; L-57 to A-71; S-58 to L-72; C-59 to Q-73; C-60 to G-74; L-61 to D-75; T-62 to L-76; V-63 to A-77; V-64 to S-78; S-65 to L-79; F-66 to R-80; Y-67 to A-81; O-68 to E-82; V-69 to L-83; A-70 to O-84; A-71 to G-85; L-72 to H-86; Q-73 to H-87; G-74 to A-88; D-75 to E-89; L-76 to K-90; A-77 to L-91; S-78 to P-92; L-79 to

A-93; R-80 to G-94; A-81 to A-95; E-82 to G-96; L-83 to A-97; Q-84 to P-98; G-85 to K-99; H-86 to A-100; H-87 to G-101; A-88 to L-102; E-89 to E-103; K-90 to E-104; L-91 to A-105; P-92 to P-106; A-93 to A-107; G-94 to V-108; A-95 to T-109; G-96 to A-110; A-97 to G-111; P-98 to L-112; K-99 to K-113; A-100 to I-114; G-101 to F-115; L-102 to E-116; E-103 to P-117; E-104 to P-118; A-105 to A-119; P-106 to P-120; A-107 to G-121; V-108 to E-122; T-109 to G-123; A-110 to N-124; G-111 to S-125; L-112 to S-126; K-113 to Q-127; I-114 to N-128; F-115 to S-129; E-116 to R-130; P-117 to N-131; P-118 to K-132; A-119 to R-133; P-120 to A-134; G-121 to V-135; E-122 to Q-136; G-123 to G-137; N-124 to P-138; S-125 to E-139; S-126 to E-140; Q-127 to T-141; N-128 to V-142; S-129 to T-143; R-130 to Q-144; N-131 to D-145; K-132 to C-146; R-133 to L-147; A-134 to Q-148; V-135 to L-149; Q-136 to I-150; G-137 to A-151; P-138 to D-152; E-139 to S-153; E-140 to E-154; T-141 to T-155; V-142 to P-156; T-143 to T-157; Q-144 to I-158; D-145 to O-159; C-146 to K-160; L-147 to G-161; Q-148 to S-162; L-149 to Y-163; I-150 to T-164; A-151 to F-165; D-152 to V-166; S-153 to P-167; E-154 to W-168; T-155 to L-169; P-156 to L-170; T-157 to S-171; I-158 to F-172; Q-159 to K-173; K-160 to R-174; G-161 to G-175; S-162 to S-176; Y-163 to A-177; T-164 to L-178; F-165 to E-179; V-166 to E-180; P-167 to K-181; W-168 to E-182; L-169 to N-183; L-170 to K-184; S-171 to I-185; F-172 to L-186; K-173 to V-187; R-174 to K-188; G-175 to E-189; S-176 to T-190; A-177 to G-191; L-178 to Y-192; E-179 to F-193; E-180 to F-194; K-181 to I-195; E-182 to Y-196; N-183 to G-197; K-184 to Q-198; I-185 to V-199; L-186 to L-200; V-187 to Y-201; K-188 to T-202; E-189 to D-203; T-190 to K-204; G-191 to T-205; Y-192 to Y-206; F-193 to A-207; F-194 to M-208; I-195 to G-209; Y-196 to H-210; G-197 to L-211; Q-198 to I-212; V-199 to Q-213; L-200 to R-214; Y-201 to K-215; T-202 to K-216; D-203 to V-217; K-204 to H-218; T-205 to V-219; Y-206 to F-220; A-207 to G-221; M-208 to D-222; G-209 to E-223; H-210 to L-224; L-211 to S-225; I-212 to L-226; Q-213 to V-227: R-214 to T-228; K-215 to L-229; K-216 to F-230; V-217 to R-231; H-218 to C-232; V-219 to I-233; F-220 to Q-234; G-221 to N-235; D-222 to M-236; E-223 to P-237; L-224 to E-238; S-225 to T-239; L-226 to L-240; V-227 to P-241; T-228 to N-242; L-229 to N-243; F-230 to S-244; R-231 to C-245; C-232 to Y-246; I-233 to S-247; Q-234 to A-248; N-235 to G-249; M-236 to I-250; P-237 to A-251; E-238 to K-252; T-239 to L-253; L-240 to E-254; P-241 to E-255; N-242 to G-256; N-243 to D-257; S-244 to E-258; C-245 to L-259; Y-246 to Q-260; S-247 to L-261; A-248 to A-262; G-249 to I-263; I-250 to P-264;

A-251 to R-265; K-252 to E-266; L-253 to N-267; E-254 to A-268; E-255 to Q-269; G-256 to I-270; D-257 to S-271; E-258 to L-272; L-259 to D-273; Q-260 to G-274; L-261 to D-275; A-262 to V-276; I-263 to T-277; P-264 to F-278; R-265 to F-279; E-266 to G-280; N-267 to A-281; A-268 to L-282; Q-269 to K-283; I-270 to L-284; and S-271 to L-285 of SEQ ID NO:3228. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

In additional embodiments, antibodies of the present invention may bind [0144] polypeptide fragments comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues: M-1 to C-15; D-2 to L-16; D-3 to K-17; S-4 to K-18; T-5 to R-19; E-6 to E-20; R-7 to E-21; E-8 to M-22; Q-9 to K-23; S-10 to L-24; R-11 to K-25; L-12 to E-26; T-13 to C-27; S-14 to V-28; C-15 to S-29; L-16 to I-30; K-17 to L-31; K-18 to P-32; R-19 to R-33; E-20 to K-34; E-21 to E-35; M-22 to S-36; K-23 to P-37; L-24 to S-38; K-25 to V-39; E-26 to R-40; C-27 to S-41; V-28 to S-42; S-29 to K-43; I-30 to D-44; L-31 to G-45; P-32 to K-46; R-33 to L-47; K-34 to L-48; E-35 to A-49; S-36 to A-50; P-37 to T-51; S-38 to L-52; V-39 to L-53; R-40 to L-54; S-41 to A-55; S-42 to L-56; K-43 to L-57; D-44 to S-58; G-45 to C-59; K-46 to C-60; L-47 to L-61; L-48 to T-62; A-49 to V-63; A-50 to V-64; T-51 to S-65; L-52 to F-66; L-53 to Y-67; L-54 to Q-68; A-55 to V-69; L-56 to A-70; L-57 to A-71; S-58 to L-72; C-59 to Q-73; C-60 to G-74; L-61 to D-75; T-62 to L-76; V-63 to A-77; V-64 to S-78; S-65 to L-79; F-66 to R-80; Y-67 to A-81; Q-68 to E-82; V-69 to L-83; A-70 to Q-84; A-71 to G-85; L-72 to H-86; Q-73 to H-87; G-74 to A-88; D-75 to E-89; L-76 to K-90; A-77 to L-91; S-78 to P-92; L-79 to A-93; R-80 to G-94; A-81 to A-95; E-82 to G-96; L-83 to A-97; Q-84 to P-98; G-85 to K-99; H-86 to A-100; H-87 to G-101; A-88 to L-102; E-89 to E-103; K-90 to E-104; L-91 to A-105; P-92 to P-106; A-93 to A-107; G-94 to V-108; A-95 to T-109; G-96 to A-110; A-97 to G-111; P-98 to L-112; K-99 to K-113; A-100 to I-114; G-101 to F-115; L-102 to E-116; E-103 to P-117; E-104 to P-118; A-105 to A-119; P-106 to P-120; A-107 to G-121; V-108 to E-122; T-109 to G-123; A-110 to N-124; G-111 to S-125; L-112 to S-126; K-113 to Q-127; I-114 to N-128; F-115 to S-129; E-116 to R-130; P-117 to N-131; P-118 to K-132; A-119 to R-133; P-120 to A-134; G-121 to V-135; E-122 to Q-136; G-123 to G-137; N-124 to P-138; S-125 to E-139; S-126 to E-140; Q-127 to T-141; N-128 to G-142;

S-129 to S-143; R-130 to Y-144; N-131 to T-145; K-132 to F-146; R-133 to V-147; A-134 to P-148; V-135 to W-149; O-136 to L-150; G-137 to L-151; P-138 to S-152; E-139 to F-153; E-140 to K-154; T-141 to R-155; G-142 to G-156; S-143 to S-157; Y-144 to A-158; T-145 to L-159; F-146 to E-160; V-147 to E-161; P-148 to K-162; W-149 to E-163; L-150 to N-164; L-151 to K-165; S-152 to I-166; F-153 to L-167; K-154 to V-168; R-155 to K-169; G-156 to E-170; S-157 to T-171; A-158 to G-172; L-159 to Y-173; E-160 to F-174; E-161 to F-175; K-162 to I-176; E-163 to Y-177; N-164 to G-178; K-165 to O-179; I-166 to V-180; L-167 to L-181; V-168 to Y-182; K-169 to T-183; E-170 to D-184; T-171 to K-185; G-172 to T-186; Y-173 to Y-187; F-174 to A-188; F-175 to M-189; I-176 to G-190: Y-177 to H-191: G-178 to L-192; O-179 to I-193; V-180 to Q-194; L-181 to R-195; Y-182 to K-196; T-183 to K-197; D-184 to V-198; K-185 to H-199; T-186 to V-200; Y-187 to F-201; A-188 to G-202; M-189 to D-203; G-190 to E-204; H-191 to L-205; L-192 to S-206; I-193 to L-207; Q-194 to V-208; R-195 to T-209; K-196 to L-210; K-197 to F-211; V-198 to R-212; H-199 to C-213; V-200 to I-214; F-201 to Q-215; G-202 to N-216; D-203 to M-217; E-204 to P-218; L-205 to E-219; S-206 to T-220; L-207 to L-221; V-208 to P-222; T-209 to N-223; L-210 to N-224; F-211 to S-225; R-212 to C-226; C-213 to Y-227: I-214 to S-228; O-215 to A-229; N-216 to G-230; M-217 to I-231; P-218 to A-232; E-219 to K-233; T-220 to L-234; L-221 to E-235; P-222 to E-236; N-223 to G-237; N-224 to D-238; S-225 to E-239; C-226 to L-240; Y-227 to Q-241; S-228 to L-242; A-229 to A-243; G-230 to I-244; I-231 to P-245; A-232 to R-246; K-233 to E-247; L-234 to N-248; E-235 to A-249; E-236 to Q-250; G-237 to I-251; D-238 to S-252; E-239 to L-253; L-240 to D-254; O-241 to G-255; L-242 to D-256; A-243 to V-257; I-244 to T-258; P-245 to F-259; R-246 to F-260; E-247 to G-261; N-248 to A-262; A-249 to L-263; Q-250 to K-264; I-251 to L-265; and S-252 to L-266 of SEQ ID NO:3229. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0145] In additional embodiments, antibodies of the present invention may bind polypeptide fragments comprising, or alternatively consisting of, an amino acid sequence selected from the group consisting of residues: M-1 to F-15; D-2 to C-16; E-3 to S-17; S-4 to E-18; A-5 to K-19; K-6 to G-20; T-7 to E-21; L-8 to D-22; P-9 to M-23; P-10 to K-24;

P-11 to V-25; C-12 to G-26; L-13 to Y-27; C-14 to D-28; F-15 to P-29; C-16 to I-30; S-17 to T-31; E-18 to P-32; K-19 to Q-33; G-20 to K-34; E-21 to E-35; D-22 to E-36; M-23 to G-37; K-24 to A-38; V-25 to W-39; G-26 to F-40; Y-27 to G-41; D-28 to I-42; P-29 to C-43; I-30 to R-44; T-31 to D-45; P-32 to G-46; Q-33 to R-47; K-34 to L-48; E-35 to L-49; E-36 to A-50; G-37 to A-51; A-38 to T-52; W-39 to L-53; F-40 to L-54; G-41 to L-55; I-42 to A-56; C-43 to L-57; R-44 to L-58; D-45 to S-59; G-46 to S-60; R-47 to S-61; L-48 to F-62; L-49 to T-63; A-50 to A-64; A-51 to M-65; T-52 to S-66; L-53 to L-67; L-54 to Y-68; L-55 to Q-69; A-56 to L-70; L-57 to A-71; L-58 to A-72; S-59 to L-73; S-60 to Q-74; S-61 to A-75; F-62 to D-76; T-63 to L-77; A-64 to M-78; M-65 to N-79; S-66 to L-80; L-67 to R-81; Y-68 to M-82; Q-69 to E-83; L-70 to L-84; A-71 to Q-85; A-72 to S-86; L-73 to Y-87; Q-74 to R-88; A-75 to G-89; D-76 to S-90; L-77 to A-91; M-78 to T-92; N-79 to P-93; L-80 to A-94; R-81 to A-95; M-82 to A-96; E-83 to G-97; L-84 to A-98; O-85 to P-99; S-86 to E-100; Y-87 to L-101; R-88 to T-102; G-89 to A-103; S-90 to G-104; A-91 to V-105; T-92 to K-106; P-93 to L-107; A-94 to L-108; A-95 to T-109; A-96 to P-110; G-97 to A-111; A-98 to A-112; P-99 to P-113; E-100 to R-114; L-101 to P-115; T-102 to H-116; A-103 to N-117; G-104 to S-118; V-105 to S-119; K-106 to R-120; L-107 to G-121; L-108 to H-122; T-109 to R-123; P-110 to N-124; A-111 to R-125; A-112 to R-126; P-113 to A-127; R-114 to F-128; P-115 to Q-129; H-116 to G-130; N-117 to P-131; S-118 to E-132; S-119 to E-133; R-120 to T-134; G-121 to E-135; H-122 to O-136; R-123 to D-137; N-124 to V-138; R-125 to D-139; R-126 to L-140; A-127 to S-141; F-128 to A-142; Q-129 to P-143; G-130 to P-144; P-131 to A-145; E-132 to P-146; E-133 to C-147; T-134 to L-148; E-135 to P-149; Q-136 to G-150; D-137 to C-151; V-138 to R-152; D-139 to H-153; L-140 to S-154; S-141 to Q-155; A-142 to H-156; P-143 to D-157; P-144 to D-158; A-145 to N-159; P-146 to G-160; C-147 to M-161; L-148 to N-162; P-149 to L-163; G-150 to R-164; C-151 to N-165; R-152 to I-166; H-153 to I-167; S-154 to Q-168; Q-155 to D-169; H-156 to C-170; D-157 to L-171; D-158 to O-172; N-159 to L-173; G-160 to I-174; M-161 to A-175; N-162 to D-176; L-163 to S-177; R-164 to D-178; N-165 to T-179; I-166 to P-180; I-167 to A-181; Q-168 to L-182; D-169 to E-183; C-170 to E-184; L-171 to K-185; Q-172 to E-186; L-173 to N-187; I-174 to K-188; A-175 to I-189; D-176 to V-190; S-177 to V-191; D-178 to R-192; T-179 to Q-193; P-180 to T-194; A-181 to G-195; L-182 to Y-196; E-183 to F-197; E-184 to F-198; K-185 to I-199; E-186 to Y-200; N-187 to S-201; K-188 to Q-202; I-189 to V-203; V-190 to L-204; V-191 to Y-205; R-192 to T-

206; Q-193 to D-207; T-194 to P-208; G-195 to I-209; Y-196 to F-210; F-197 to A-211; F-198 to M-212; I-199 to G-213; Y-200 to H-214; S-201 to V-215; Q-202 to I-216; V-203 to Q-217; L-204 to R-218; Y-205 to K-219; T-206 to K-220; D-207 to V-221; P-208 to H-222; I-209 to V-223; F-210 to F-224; A-211 to G-225; M-212 to D-226; G-213 to E-227; H-214 to L-228; V-215 to S-229; I-216 to L-230; Q-217 to V-231; R-218 to T-232; K-219 to L-233; K-220 to F-234; V-221 to R-235; H-222 to C-236; V-223 to I-237; F-224 to Q-238; G-225 to N-239; D-226 to M-240; E-227 to P-241; L-228 to K-242; S-229 to T-243; L-230 to L-244; V-231 to P-245; T-232 to N-246; L-233 to N-247; F-234 to S-248; R-235 to C-249; C-236 to Y-250; I-237 to S-251; Q-238 to A-252; N-239 to G-253; M-240 to I-254; P-241 to A-255; K-242 to R-256; T-243 to L-257; L-244 to E-258; P-245 to E-259; N-246 to G-260; N-247 to D-261; S-248 to E-262; C-249 to I-263; Y-250 to Q-264; S-251 to L-265; A-252 to A-266; G-253 to I-267; I-254 to P-268; A-255 to R-269; R-256 to E-270; L-257 to N-271; E-258 to A-272; E-259 to Q-273; G-260 to I-274; D-261 to S-275; E-262 to R-276; I-263 to N-277; Q-264 to G-278; L-265 to D-279; A-266 to D-280; I-267 to T-281; P-268 to F-282; R-269 to F-283; E-270 to G-284; N-271 to A-285; A-272 to L-286; O-273 to K-287; I-274 to L-288; and S-275 to L-289 of SEQ ID NO:38. The present invention is also directed to antibodies that bind BLyS polypeptides comprising, or alternatively, consisting of, a contiguous sequence of amino acid residues at least 80%, 85%, 90%, 92%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of BLyS polypeptides described above.

[0146] It will be recognized by one of ordinary skill in the art that some amino acid sequences of the BLyS polypeptides can be varied without significant effect of the structure or function of the polypeptide. If such differences in sequence are contemplated, it should be remembered that there will be critical areas on the polypeptide which determine activity.

[0147] Thus, the invention further includes antibodies that bind variations of BLyS polypeptides which show BLyS polypeptide functional activity (e.g., biological activity) or which include regions of BLyS polypeptide such as the polypeptide fragments described herein. Such mutants include deletions, insertions, inversions, repeats, and type substitutions selected according to general rules known in the art so as have little effect on activity. For example, guidance concerning how to make phenotypically silent amino acid substitutions is provided in Bowie, J. U. et al., "Deciphering the Message in Protein

Sequences: Tolerance to Amino Acid Substitutions," Science 247:1306-1310 (1990), wherein the authors indicate that there are two main approaches for studying the tolerance of an amino acid sequence to change. The first method relies on the process of evolution, in which mutations are either accepted or rejected by natural selection. The second approach uses genetic engineering to introduce amino acid changes at specific positions of a cloned gene and selections or screens to identify sequences that maintain functionality.

[0148] As the authors state, these studies have revealed that proteins are surprisingly tolerant of amino acid substitutions. The authors further indicate which amino acid changes are likely to be permissive at a certain position of the protein. For example, most buried amino acid residues require nonpolar side chains, whereas few features of surface side chains are generally conserved. Other such phenotypically silent substitutions are described in Bowie, J. U. et al., supra, and the references cited therein. Typically seen as conservative substitutions are the replacements, one for another, among the aliphatic amino acids Ala, Val, Leu and Ile; interchange of the hydroxyl residues Ser and Thr, exchange of the acidic residues Asp and Glu, substitution between the amide residues Asn and Gln, exchange of the basic residues Lys and Arg and replacements among the aromatic residues Phe, Tyr.

[0149] Thus, antibodies of the present invention may bind fragments, derivatives or analogs of the polypeptide of SEQ ID NO:3228, or that encoded by the deposited cDNA plasmid, such as (i) polypeptides in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue) and such substituted amino acid residue may or may not be one encoded by the genetic code, or (ii) polypeptides in which one or more of the amino acid residues includes a substituent group, or (iii) polypeptides in which the extracellular domain of the polypeptide is fused with another compound, such as a compound to increase the half-life of the polypeptide (for example, polyethylene glycol), or (iv) polypeptides in which the additional amino acids are fused to the extracellular domain of the polypeptide, such as an IgG Fc fusion region peptide or leader or secretory sequence or a sequence which is employed for purification of the extracellular domain of the polypeptide or a proprotein sequence.

[0150] Antibodies of the present invention may bind fragments, derivatives or analogs of the polypeptide of SEQ ID NO:3229, or that encoded by the deposited cDNA

plasmid, such as (i) polypeptides in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue) and such substituted amino acid residue may or may not be one encoded by the genetic code, or (ii) polypeptides in which one or more of the amino acid residues includes a substituent group, or (iii) polypeptides in which the extracellular domain of the polypeptide is fused with another compound, such as a compound to increase the half-life of the polypeptide (for example, polyethylene glycol), or (iv) polypeptides in which the additional amino acids are fused to the extracellular domain of the polypeptide, such as, a soluble biologically active fragment of another TNF ligand family member (e.g., CD40 Ligand), an IgG Fc fusion region peptide or leader or secretory sequence or a sequence which is employed for purification of the extracellular domain of the polypeptide or a proprotein sequence. Such fragments, derivatives and analogs are deemed to be within the scope of those skilled in the art from the teachings herein.

[0151] Thus, the antibodies of the invention may bind BLyS polypeptides that include one or more amino acid substitutions, deletions or additions, either from natural mutations or human manipulation. As indicated, changes are preferably of a minor nature, such as conservative amino acid substitutions that do not significantly affect the folding or activity of the protein (see Table 13).

[0152	TABLE 13. Conservative Amino Acid Substitutions.					
	[0153]	Aromatic	[0159]	Phenylalanine		
			[0160]	Tryptophan	1	
			[0161]	Tyrosine		
1	[0154]	Hydrophobic	[0162]	Leucine		
		· •	[0163]	Isoleucine		
			[0164]	Valine		
	[0155]	Polar	[0165]	Glutamine	-	
			[0166]	Asparagine		
	[0156]	Basic	[0167]	Arginine	- 1	
	•		101681	Lysine	i	

[0157]

Acidic

[0169]

[0170]

[0171]

Histidine

Aspartic Acid

Glutamic Acid

	_ ••				
[0158]	Small		[0172]	Alanine	
		· I	[0173]	Serine	
			[0174]	Threonine	
]		Ì	[0175]	Methionine	
			[0176]	Glycine	

In one embodiment of the invention, antibodies of the present invention

[0177]

bind polypeptides comprising, or alternatively consisting of, the amino acid sequence of a BLyS polypeptide having an amino acid sequence which contains at least one conservative amino acid substitution, but not more than 50 conservative amino acid substitutions, even more preferably, not more than 40 conservative amino acid substitutions, still more preferably, not more than 30 conservative amino acid substitutions, and still even more preferably, not more than 20 conservative amino acid substitutions. In one embodiment of the invention, antibodies of the present invention bind polypeptides comprising, or alternatively consisting of, the amino acid sequence of a BLyS polypeptide having an amino acid sequence which contains at least one conservative amino acid substitution, but not more than 10, 9, 8, 7, 6, 5, 4, 3, 2 or 1 conservative amino acid substitutions. For example, site directed changes at the amino acid level of BLyS can be [0178] made by replacing a particular amino acid with a conservative substitution. Antibodies of the present invention may bind BLyS amino acid sequences containing conservative substitution mutations of the polypeptide of SEQ ID NO:3228 including: M1 replaced with A, G, I, L, S, T, or V; D2 replaced with E; D3 replaced with E; S4 replaced with A, G. I. L. T. M. or V: T5 replaced with A. G. I. L. S. M. or V: E6 replaced with D; R7 replaced with H, or K; E8 replaced with D; Q9 replaced with N; S10 replaced with A, G, I, L, T, M, or V; R11 replaced with H, or K; L12 replaced with A, G, I, S, T, M, or V; T13 replaced with A, G, I, L, S, M, or V; S14 replaced with A, G, I, L, T, M, or V; L16 replaced with A, G, I, S, T, M, or V; K17 replaced with H, or R; K18 replaced with H, or R; R19 replaced with H, or K; E20 replaced with D; E21 replaced with D; M22 replaced with A, G, I, L, S, T, or V; K23 replaced with H, or R; L24 replaced with A, G, I, S, T, M, or V; K25 replaced with H, or R; E26 replaced with D; V28 replaced with A, G, I, L, S, T, or M; S29 replaced with A, G, I, L, T, M, or V; I30 replaced with A, G, L, S, T, M, or V; L31 replaced with A, G, I, S, T, M, or V; R33 replaced with H, or K; K34 replaced with H, or R; E35 replaced with D; S36 replaced with A, G, I, L, T, M, or V; S38 replaced with

A, G, I, L, T, M, or V; V39 replaced with A, G, I, L, S, T, or M; R40 replaced with H, or K; S41 replaced with A, G, I, L, T, M, or V; S42 replaced with A, G, I, L, T, M, or V; K43 replaced with H, or R; D44 replaced with E; G45 replaced with A, I, L, S, T, M, or V; K46 replaced with H, or R; L47 replaced with A, G, I, S, T, M, or V; L48 replaced with A, G, I, S, T, M, or V; A49 replaced with G, I, L, S, T, M, or V; A50 replaced with G, I, L, S, T, M, or V; T51 replaced with A, G, I, L, S, M, or V; L52 replaced with A, G, I, S, T, M, or V; L53 replaced with A, G, I, S, T, M, or V; L54 replaced with A, G, I, S, T, M, or V; A55 replaced with G, I, L, S, T, M, or V; L56 replaced with A, G, I, S, T, M, or V; L57 replaced with A, G, I, S, T, M, or V; S58 replaced with A, G, I, L, T, M, or V; L61 replaced with A, G, I, S, T, M, or V; T62 replaced with A, G, I, L, S, M, or V; V63 replaced with A, G, I, L, S, T, or M; V64 replaced with A, G, I, L, S, T, or M; S65 replaced with A, G, I, L, T, M, or V; F66 replaced with W, or Y; Y67 replaced with F, or W; O68 replaced with N; V69 replaced with A, G, I, L, S, T, or M; A70 replaced with G, I, L, S, T, M, or V; A71 replaced with G, I, L, S, T, M, or V; L72 replaced with A, G, I, S, T. M. or V: O73 replaced with N; G74 replaced with A, I, L, S, T, M, or V; D75 replaced with E; L76 replaced with A, G, I, S, T, M, or V; A77 replaced with G, I, L, S, T, M, or V; S78 replaced with A, G, I, L, T, M, or V; L79 replaced with A, G, I, S, T, M, or V; R80 replaced with H, or K; A81 replaced with G, I, L, S, T, M, or V; E82 replaced with D; L83 replaced with A, G, I, S, T, M, or V; Q84 replaced with N; G85 replaced with A, I, L, S, T, M, or V; H86 replaced with K, or R; H87 replaced with K, or R; A88 replaced with G, I, L, S, T, M, or V; E89 replaced with D; K90 replaced with H, or R; L91 replaced with A, G, I, S, T, M, or V; A93 replaced with G, I, L, S, T, M, or V; G94 replaced with A, I, L, S, T, M, or V; A95 replaced with G, I, L, S, T, M, or V; G96 replaced with A, I, L, S, T, M, or V; A97 replaced with G, I, L, S, T, M, or V; K99 replaced with H, or R; A100 replaced with G, I, L, S, T, M, or V; G101 replaced with A, I, L, S, T, M, or V; L102 replaced with A, G, I, S, T, M, or V; E103 replaced with D; E104 replaced with D; A105 replaced with G, I, L, S, T, M, or V; A107 replaced with G, I, L, S, T, M, or V; V108 replaced with A, G, I, L, S, T, or M; T109 replaced with A, G, I, L, S, M, or V; A110 replaced with G, I, L, S, T, M, or V; G111 replaced with A, I, L, S, T, M, or V; L112 replaced with A, G, I, S, T, M, or V; K113 replaced with H, or R; I114 replaced with A, G, L, S, T, M, or V; F115 replaced with W, or Y; E116 replaced with D; A119 replaced with G, I, L, S, T, M, or V; G121 replaced with A, I, L, S, T, M, or V; E122 replaced with D; G123 replaced with A,

I, L, S, T, M, or V; N124 replaced with Q; S125 replaced with A, G, I, L, T, M, or V; S126 replaced with A, G, I, L, T, M, or V; Q127 replaced with N; N128 replaced with Q; S129 replaced with A, G, I, L, T, M, or V; R130 replaced with H, or K; N131 replaced with Q; K132 replaced with H, or R; R133 replaced with H, or K; A134 replaced with G. I, L, S, T, M, or V; V135 replaced with A, G, I, L, S, T, or M; Q136 replaced with N; G137 replaced with A, I, L, S, T, M, or V; E139 replaced with D; E140 replaced with D: T141 replaced with A, G, I, L, S, M, or V; V142 replaced with A, G, I, L, S, T, or M; T143 replaced with A, G, I, L, S, M, or V; O144 replaced with N; D145 replaced with E; L147 replaced with A, G, I, S, T, M, or V; Q148 replaced with N; L149 replaced with A, G, I, S, T, M, or V; I150 replaced with A, G, L, S, T, M, or V; A151 replaced with G, I, L, S, T, M, or V; D152 replaced with E; S153 replaced with A, G, I, L, T, M, or V: E154 replaced with D; T155 replaced with A, G, I, L, S, M, or V; T157 replaced with A, G, I, L. S, M, or V; I158 replaced with A, G, L, S, T, M, or V; Q159 replaced with N; K160 replaced with H, or R; G161 replaced with A, I, L, S, T, M, or V; S162 replaced with A, G, I, L, T, M, or V; Y163 replaced with F, or W; T164 replaced with A, G, I, L, S, M, or V; F165 replaced with W, or Y; V166 replaced with A, G, I, L, S, T, or M; W168 replaced with F, or Y; L169 replaced with A, G, I, S, T, M, or V; L170 replaced with A, G, I, S, T, M, or V; S171 replaced with A, G, I, L, T, M, or V; F172 replaced with W, or Y; K173 replaced with H, or R; R174 replaced with H, or K; G175 replaced with A, I, L, S, T, M, or V; S176 replaced with A, G, I, L, T, M, or V; A177 replaced with G, I, L, S, T, M, or V; L178 replaced with A, G, I, S, T, M, or V; E179 replaced with D; E180 replaced with D; K181 replaced with H, or R; E182 replaced with D; N183 replaced with O; K184 replaced with H, or R; I185 replaced with A, G, L, S, T, M, or V; L186 replaced with A, G, I, S, T, M, or V; V187 replaced with A, G, I, L, S, T, or M; K188 replaced with H, or R; E189 replaced with D; T190 replaced with A, G, I, L, S, M, or V; G191 replaced with A, I, L, S, T, M, or V; Y192 replaced with F, or W; F193 replaced with W, or Y; F194 replaced with W, or Y; I195 replaced with A, G, L, S, T, M, or V; Y196 replaced with F, or W; G197 replaced with A, I, L, S, T, M, or V; Q198 replaced with N; V199 replaced with A, G, I, L, S, T, or M; L200 replaced with A, G, I, S, T, M, or V; Y201 replaced with F, or W; T202 replaced with A, G, I, L, S, M, or V; D203 replaced with E; K204 replaced with H, or R; T205 replaced with A, G, I, L, S, M, or V; Y206 replaced with F, or W; A207 replaced with G, I, L, S, T, M, or V; M208 replaced with A, G, I, L, S, T, or V;

G209 replaced with A, I, L, S, T, M, or V; H210 replaced with K, or R; L211 replaced with A, G, I, S, T, M, or V; I212 replaced with A, G, L, S, T, M, or V; Q213 replaced with N; R214 replaced with H, or K; K215 replaced with H, or R; K216 replaced with H, or R; V217 replaced with A, G, I, L, S, T, or M; H218 replaced with K, or R; V219 replaced with A, G, I, L, S, T, or M; F220 replaced with W, or Y; G221 replaced with A, I, L, S, T, M, or V; D222 replaced with E; E223 replaced with D; L224 replaced with A, G, I, S, T, M, or V; S225 replaced with A, G, I, L, T, M, or V; L226 replaced with A, G, I, S, T, M. or V; V227 replaced with A, G, I, L, S, T, or M; T228 replaced with A, G, I, L, S, M, or V; L229 replaced with A, G, I, S, T, M, or V; F230 replaced with W, or Y; R231 replaced with H, or K; I233 replaced with A, G, L, S, T, M, or V; Q234 replaced with N; N235 replaced with Q; M236 replaced with A, G, I, L, S, T, or V; E238 replaced with D; T239 replaced with A, G, I, L, S, M, or V; L240 replaced with A, G, I, S, T, M, or V; N242 replaced with Q; N243 replaced with Q; S244 replaced with A, G, I, L, T, M, or V; Y246 replaced with F, or W; S247 replaced with A, G, I, L, T, M, or V; A248 replaced with G, I, L, S, T, M, or V; G249 replaced with A, I, L, S, T, M, or V; I250 replaced with A, G, L, S, T, M, or V; A251 replaced with G, I, L, S, T, M, or V; K252 replaced with H, or R; L253 replaced with A, G, I, S, T, M, or V; E254 replaced with D; E255 replaced with D; G256 replaced with A, I, L, S, T, M, or V; D257 replaced with E; E258 replaced with D: L259 replaced with A, G, I, S, T, M, or V; Q260 replaced with N; L261 replaced with A, G, I, S, T, M, or V; A262 replaced with G, I, L, S, T, M, or V; I263 replaced with A, G, L, S, T, M, or V; R265 replaced with H, or K; E266 replaced with D; N267 replaced with Q; A268 replaced with G, I, L, S, T, M, or V; Q269 replaced with N; I270 replaced with A, G, L, S, T, M, or V; S271 replaced with A, G, I, L, T, M, or V; L272 replaced with A, G, I, S, T, M, or V; D273 replaced with E; G274 replaced with A, I, L, S, T, M, or V; D275 replaced with E; V276 replaced with A, G, I, L, S, T, or M; T277 replaced with A, G, I, L, S. M. or V; F278 replaced with W, or Y; F279 replaced with W, or Y; G280 replaced with A, I, L, S, T, M, or V; A281 replaced with G, I, L, S, T, M, or V; L282 replaced with A, G, I, S, T, M, or V; K283 replaced with H, or R; L284 replaced with A, G, I, S, T, M, or V; and/or L285 replaced with A, G, I, S, T, M, or V.

[0179] In another embodiment, site directed changes at the amino acid level of BLyS can be made by replacing a particular amino acid with a conservative substitution. Antibodies of the present invention may bind BLyS amino acid sequences containing

conservative substitution mutations of the polypeptide of SEQ ID NO:3229 including: M1 replaced with A, G, I, L, S, T, or V; D2 replaced with E; D3 replaced with E; S4 replaced with A, G, I, L, T, M, or V; T5 replaced with A, G, I, L, S, M, or V; E6 replaced with D; R7 replaced with H, or K; E8 replaced with D; Q9 replaced with N; S10 replaced with A, G, I, L, T, M, or V; R11 replaced with H, or K; L12 replaced with A, G, I, S, T, M, or V; T13 replaced with A, G, I, L, S, M, or V; S14 replaced with A, G, I, L, T, M, or V; L16 replaced with A, G, I, S, T, M, or V; K17 replaced with H, or R; K18 replaced with H, or R; R19 replaced with H, or K; E20 replaced with D; E21 replaced with D; M22 replaced with A, G, I, L, S, T, or V; K23 replaced with H, or R; L24 replaced with A, G, I, S, T, M. or V; K25 replaced with H, or R; E26 replaced with D; V28 replaced with A, G, I, L, S, T, or M; S29 replaced with A, G, I, L, T, M, or V; I30 replaced with A, G, L, S, T, M, or V; L31 replaced with A, G, I, S, T, M, or V; R33 replaced with H, or K; K34 replaced with H, or R; E35 replaced with D; S36 replaced with A, G, I, L, T, M, or V; S38 replaced with A, G, I, L, T, M, or V; V39 replaced with A, G, I, L, S, T, or M; R40 replaced with H, or K; S41 replaced with A, G, I, L, T, M, or V; S42 replaced with A, G, I, L, T, M, or V; K43 replaced with H, or R; D44 replaced with E; G45 replaced with A, I, L, S, T, M, or V; K46 replaced with H, or R; L47 replaced with A, G, I, S, T, M, or V; L48 replaced with A, G, I. S. T. M. or V: A49 replaced with G, I, L, S, T, M, or V; A50 replaced with G, I, L, S, T, M, or V; T51 replaced with A, G, I, L, S, M, or V; L52 replaced with A, G, I, S, T, M, or V; L53 replaced with A, G, I, S, T, M, or V; L54 replaced with A, G, I, S, T, M, or V; A55 replaced with G, I, L, S, T, M, or V; L56 replaced with A, G, I, S, T, M, or V; L57 replaced with A, G, I, S, T, M, or V; S58 replaced with A, G, I, L, T, M, or V; L61 replaced with A, G, I, S, T, M, or V; T62 replaced with A, G, I, L, S, M, or V; V63 replaced with A, G, I, L, S, T, or M; V64 replaced with A, G, I, L, S, T, or M; S65 replaced with A, G, I, L, T, M, or V; F66 replaced with W, or Y; Y67 replaced with F, or W; Q68 replaced with N; V69 replaced with A, G, I, L, S, T, or M; A70 replaced with G. I, L, S, T, M, or V; A71 replaced with G, I, L, S, T, M, or V; L72 replaced with A, G, I, S, T, M, or V; Q73 replaced with N; G74 replaced with A, I, L, S, T, M, or V; D75 replaced with E; L76 replaced with A, G, I, S, T, M, or V; A77 replaced with G, I, L, S, T, M, or V; S78 replaced with A, G, I, L, T, M, or V; L79 replaced with A, G, I, S, T, M, or V; R80 replaced with H, or K; A81 replaced with G, I, L, S, T, M, or V; E82 replaced with D; L83 replaced with A, G, I, S, T, M, or V; Q84 replaced with N; G85 replaced with A, I, L, S,

T, M, or V; H86 replaced with K, or R; H87 replaced with K, or R; A88 replaced with G, I, L, S, T, M, or V; E89 replaced with D; K90 replaced with H, or R; L91 replaced with A, G, I, S, T, M, or V; A93 replaced with G, I, L, S, T, M, or V; G94 replaced with A, I, L, S, T, M, or V; A95 replaced with G, I, L, S, T, M, or V; G96 replaced with A, I, L, S, T, M, or V; A97 replaced with G, I, L, S, T, M, or V; K99 replaced with H, or R; A100 replaced with G, I, L, S, T, M, or V; G101 replaced with A, I, L, S, T, M, or V; L102 replaced with A, G, I, S, T, M, or V; E103 replaced with D; E104 replaced with D; A105 replaced with G, I, L, S, T, M, or V; A107 replaced with G, I, L, S, T, M, or V; V108 replaced with A, G, I, L, S, T, or M; T109 replaced with A, G, I, L, S, M, or V; A110 replaced with G, I, L, S, T, M, or V; G111 replaced with A, I, L, S, T, M, or V; L112 replaced with A, G, I, S, T, M, or V; K113 replaced with H, or R; I114 replaced with A, G, L, S, T, M, or V; F115 replaced with W, or Y; E116 replaced with D; A119 replaced with G, I, L, S, T, M, or V; G121 replaced with A, I, L, S, T, M, or V; E122 replaced with D; G123 replaced with A, I, L, S, T, M, or V; N124 replaced with Q; S125 replaced with A, G, I, L, T, M, or V; S126 replaced with A, G, I, L, T, M, or V; Q127 replaced with N; N128 replaced with Q; S129 replaced with A, G, I, L, T, M, or V; R130 replaced with H, or K; N131 replaced with Q; K132 replaced with H, or R; R133 replaced with H, or K; A134 replaced with G, I, L, S, T, M, or V; V135 replaced with A, G, I, L, S, T, or M; Q136 replaced with N; G137 replaced with A, I, L, S, T, M, or V; E139 replaced with D; E140 replaced with D; T141 replaced with A, G, I, L, S, M, or V; G142 replaced with A, I, L, S, T, M, or V; S143 replaced with A, G, I, L, T, M, or V; Y144 replaced with F, or W; T145 replaced with A, G, I, L, S, M, or V; F146 replaced with W, or Y; V147 replaced with A, G, I, L, S, T, or M; W149 replaced with F, or Y; L150 replaced with A, G, I, S, T, M, or V; L151 replaced with A, G, I, S, T, M, or V; S152 replaced with A, G, I, L, T, M, or V; F153 replaced with W, or Y; K154 replaced with H, or R; R155 replaced with H, or K; G156 replaced with A, I, L, S, T, M, or V; S157 replaced with A, G, I, L, T, M, or V; A158 replaced with G, I, L, S, T, M, or V; L159 replaced with A, G, I, S, T, M, or V; E160 replaced with D; E161 replaced with D; K162 replaced with H, or R; E163 replaced with D; N164 replaced with Q; K165 replaced with H, or R; I166 replaced with A, G, L, S, T, M, or V; L167 replaced with A, G, I, S, T, M, or V; V168 replaced with A, G, I, L, S, T, or M; K169 replaced with H, or R; E170 replaced with D; T171 replaced with A, G, I, L, S, M, or V; G172 replaced with A, I, L, S, T, M, or V; Y173 replaced with F, or W; F174

replaced with W, or Y; F175 replaced with W, or Y; I176 replaced with A, G, L, S, T, M, or V; Y177 replaced with F, or W; G178 replaced with A, I, L, S, T, M, or V; Q179 replaced with N; V180 replaced with A, G, I, L, S, T, or M; L181 replaced with A, G, I, S, T, M, or V; Y182 replaced with F, or W; T183 replaced with A, G, I, L, S, M, or V; D184 replaced with E; K185 replaced with H, or R; T186 replaced with A, G, I, L, S, M, or V; Y187 replaced with F, or W; A188 replaced with G, I, L, S, T, M, or V; M189 replaced with A, G, I, L, S, T, or V; G190 replaced with A, I, L, S, T, M, or V; H191 replaced with K, or R; L192 replaced with A, G, I, S, T, M, or V; I193 replaced with A, G, L, S, T, M, or V; Q194 replaced with N; R195 replaced with H, or K; K196 replaced with H, or R; K197 replaced with H, or R; V198 replaced with A, G, I, L, S, T, or M; H199 replaced with K, or R; V200 replaced with A, G, I, L, S, T, or M; F201 replaced with W, or Y; G202 replaced with A, I, L, S, T, M, or V; D203 replaced with E; E204 replaced with D; L205 replaced with A, G, I, S, T, M, or V; S206 replaced with A, G, I, L, T, M, or V; L207 replaced with A, G, I, S, T, M, or V; V208 replaced with A, G, I, L, S, T, or M; T209 replaced with A, G, I, L, S, M, or V; L210 replaced with A, G, I, S, T, M, or V; F211 replaced with W, or Y; R212 replaced with H, or K; I214 replaced with A, G, L, S, T, M, or V; Q215 replaced with N; N216 replaced with Q; M217 replaced with A, G, I, L, S, T, or V; E219 replaced with D; T220 replaced with A, G, I, L, S, M, or V; L221 replaced with A, G, I, S, T, M, or V; N223 replaced with Q; N224 replaced with Q; S225 replaced with A, G, I, L, T, M, or V; Y227 replaced with F, or W; S228 replaced with A, G, I, L, T, M, or V; A229 replaced with G, I, L, S, T, M, or V; G230 replaced with A, I, L, S, T, M, or V; I231 replaced with A, G, L, S, T, M, or V; A232 replaced with G, I, L, S, T, M, or V; K233 replaced with H, or R; L234 replaced with A, G, I, S, T, M, or V; E235 replaced with D; E236 replaced with D; G237 replaced with A, I, L, S, T, M, or V; D238 replaced with E; E239 replaced with D; L240 replaced with A, G, I, S, T, M, or V; Q241 replaced with N; L242 replaced with A, G, I, S, T, M, or V; A243 replaced with G, I, L, S, T, M, or V; I244 replaced with A, G, L, S, T, M, or V; R246 replaced with H, or K; E247 replaced with D; N248 replaced with Q; A249 replaced with G, I, L, S, T, M, or V; Q250 replaced with N; I251 replaced with A, G, L, S, T, M, or V; S252 replaced with A, G, I, L, T, M, or V; L253 replaced with A, G, I, S, T, M, or V; D254 replaced with E; G255 replaced with A, I, L, S, T, M, or V; D256 replaced with E; V257 replaced with A, G, I, L, S, T, or M; T258 replaced with A, G, I, L, S, M, or V; F259 replaced with W, or Y; F260 replaced

with W, or Y; G261 replaced with A, I, L, S, T, M, or V; A262 replaced with G, I, L, S, T, M, or V; L263 replaced with A, G, I, S, T, M, or V; K264 replaced with H, or R; L265 replaced with A, G, I, S, T, M, or V; and/or L266 replaced with A, G, I, S, T, M, or V.

[0180] In another embodiment, site directed changes at the amino acid level of BLyS can be made by replacing a particular amino acid with a conservative substitution. Antibodies of the present invention may bind BLyS amino acid sequences containing

conservative substitution mutations of the polypeptide of any one of SEQ ID NOS:3230-

3237.

[0181] Amino acids in the BLyS polypeptides that are essential for function can be identified by methods known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis (Cunningham and Wells, Science 244:1081-1085 (1989)). The latter procedure introduces single alanine mutations at every residue in the molecule. The resulting mutant molecules are then tested for functional activity, such ligand binding and the ability to stimulate lymphocyte (e.g., B cell) as, for example, proliferation, differentiation, and/or activation. Accordingly, antibodies of the present invention may bind amino acids in the BLyS polypeptides that are essential for function. In preferred embodiments, antibodies of the present invention bind amino acids in the BLyS polypeptides that are essential for function and inhibit BLyS polypeptide function. In other preferred embodiments, antibodies of the present invention bind amino acids in the BLyS polypeptides that are essential for function and enhance BLyS polypeptide function.

[0182] Of special interest are substitutions of charged amino acids with other charged or neutral amino acids which may produce proteins with highly desirable improved characteristics, such as less aggregation. Aggregation may not only reduce activity but also be problematic when preparing pharmaceutical formulations, because aggregates can be immunogenic (Pinckard et al., Clin. Exp. Immunol. 2:331-340 (1967); Robbins et al., Diabetes 36: 838-845 (1987); Cleland et al., Crit. Rev. Therapeutic Drug Carrier Systems 10:307-377 (1993).

[0183] In another embodiment, the invention provides for antibodies that bind polypeptides having amino acid sequences containing non-conservative substitutions of the amino acid sequence provided in SEQ ID NO:3228. For example, non-conservative substitutions of the BLyS protein sequence provided in SEQ ID NO:3228 include: M1 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D2 replaced with H, K, R, A, G, I, L,

S, T, M, V, N, Q, F, W, Y, P, or C; D3 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; S4 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T5 replaced with D. E. H. K. R. N. Q. F. W. Y. P. or C; E6 replaced with H, K, R, A, G, I, L, S, T, M, V, N, O. F. W. Y. P. or C; R7 replaced with D. E. A. G. I. L. S. T. M. V. N. Q. F. W. Y. P. or C; E8 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; Q9 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; S10 replaced with D, E, H, K, R, N. O. F. W. Y. P. or C; R11 replaced with D. E. A. G. I. L. S. T. M. V. N. Q. F. W. Y. P. or C; L12 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T13 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S14 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; C15 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; L16 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K17 replaced with D, E, A, G, I, L, S, T, M, V, N, O, F, W, Y, P, or C; K18 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; R19 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E20 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E21 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; M22 replaced with D, E, H, K, R, N, Q, F, W; Y, P, or C; K23 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L24 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K25 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E26 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; C27 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; V28 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S29 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; I30 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L31 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P32 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; R33 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K34 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E35 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; S36 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P37 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; S38 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V39 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; R40 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; S41 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S42 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K43 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; D44 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G45 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or

C; K46 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L47 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L48 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A49 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A50 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T51 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L52 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L53 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L54 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A55 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L56 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L57 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S58 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; C59 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; C60 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; L61 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T62 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V63 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V64 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S65 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F66 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; Y67 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; Q68 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; V69 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A70 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A71 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L72 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q73 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; G74 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D75 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L76 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A77 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S78 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L79 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; R80 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A81 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E82 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L83 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q84 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; G85 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; H86 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; H87 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A88 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E89 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K90 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L91 replaced with

D. E. H. K. R. N. O. F. W. Y. P. or C; P92 replaced with D. E. H. K. R. A. G. I. L. S. T. M, V, N, Q, F, W, Y, or C; A93 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G94 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A95 replaced with D, E, H, K, R, N, O. F. W. Y. P. or C; G96 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A97 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P98 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; K99 replaced with D, E, A, G, I, L, S, T, M, V, N. O. F. W. Y. P. or C; A100 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G101 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L102 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E103 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E104 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A105 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P106 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; A107 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V108 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T109 replaced with D. E, H, K, R, N, Q, F, W, Y, P, or C; A110 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G111 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L112 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K113 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W. Y. P. or C: I114 replaced with D. E. H. K. R. N. Q. F. W. Y. P. or C; F115 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; E116 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; P117 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; P118 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; A119 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P120 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; G121 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E122 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F. W. Y. P. or C: G123 replaced with D. E. H. K. R. N. Q. F. W. Y. P. or C; N124 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; S125 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S126 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q127 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; N128 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; S129 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; R130 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; N131 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C: K132 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; R133 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A134 replaced with D,

E, H, K, R, N, Q, F, W, Y, P, or C; V135 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q136 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; G137 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P138 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; E139 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E140 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T141 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V142 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T143 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q144 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; D145 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; C146 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; L147 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q148 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; L149 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; I150 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A151 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D152 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; S153 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E154 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T155 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P156 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; T157 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; I158 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q159 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; K160 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G161 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S162 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y163 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; T164 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F165 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; V166 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P167 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; W168 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; L169 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L170 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S171 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F172 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; K173 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; R174 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G175 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S176 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A177 replaced with D, E, H, K, R, N, Q, F, W,

Y, P, or C; L178 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E179 replaced with H. K. R. A. G. I. L. S. T. M. V. N. Q. F. W. Y. P. or C; E180 replaced with H. K. R. A. G. I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K181 replaced with D, E, A, G, I, L, S, T, M, V, N. O. F. W. Y. P. or C; E182 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; N183 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; K184 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; I185 replaced with D, E. H. K. R. N. O. F. W. Y. P. or C; L186 replaced with D. E. H. K. R. N. Q. F. W. Y. P. or C: V187 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K188 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E189 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T190 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G191 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y192 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; F193 replaced with D, E, H, K, R, N, Q, A, G, I, L. S. T. M. V. P. or C. F194 replaced with D. E. H. K. R. N. Q. A. G. I. L. S. T. M. V. P. or C; I195 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y196 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; G197 replaced with D, E, H, K, R, N, Q, F. W. Y. P. or C; Q198 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; V199 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L200 replaced with D, E, H, K, R. N. O. F. W. Y. P. or C; Y201 replaced with D. E. H. K. R. N. Q. A. G. I. L. S. T. M. V. P, or C; T202 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D203 replaced with H, K. R. A. G. I. L. S. T. M. V. N. Q. F. W. Y. P. or C; K204 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T205 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y206 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; A207 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; M208 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G209 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; H210 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L211 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; 1212 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q213 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; R214 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K215 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K216 replaced with D, E, A, G, I, L, S, T, M, V, N, Q. F, W, Y, P, or C; V217 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; H218 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; V219 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F220 replaced with D, E, H, K, R, N, Q, A, G, I, L, S.

T. M. V. P. or C; G221 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D222 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E223 replaced with H. K. R. A. G. I. L. S. T. M. V. N. Q. F. W. Y. P. or C; L224 replaced with D. E. H. K. R. N, Q, F, W, Y, P, or C; S225 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L226 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V227 replaced with D, E, H, K, R, N, O, F, W, Y, P, or C; T228 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L229 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F230 replaced with D, E, H, K, R, N, O. A. G. I. L. S. T. M. V. P. or C; R231 replaced with D. E. A. G. I. L. S. T. M. V. N. O. F, W, Y, P, or C; C232 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; 1233 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q234 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; N235 replaced with D, E, H, K, R, A, G. I, L, S, T, M, V, F, W, Y, P, or C; M236 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C: P237 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; E238 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T239 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L240 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P241 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; N242 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; N243 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; S244 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; C245 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W. Y. or P; Y246 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; S247 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A248 replaced with D, E, H, K, R, N, O. F. W. Y. P. or C; G249 replaced with D. E. H. K. R. N. Q. F. W. Y. P. or C; I250 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A251 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K252 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L253 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E254 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E255 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G256 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D257 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E258 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L259 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q260 replaced with D, E, H, K, R, A, G, I, L, S, T. M, V, F, W, Y, P, or C; L261 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A262 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; 1263 replaced with D, E, H, K, R, N,

Q, F, W, Y, P, or C; P264 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; R265 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E266 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; N267 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; A268 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q269 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; 1270 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S271 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L272 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D273 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G274 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D275 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; V276 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T277 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F278 replaced with D. E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; F279 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; G280 replaced with D, E, H, K, R, N, O, F, W, Y, P, or C: A281 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L282 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K283 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L284 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; and/or L285 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C.

[0184] In an additional embodiment, antibodies of the present invention bind BLyS polypeptides comprising, or alternatively consisting of, a BLyS amino acid sequence in which more than one amino acid (e.g., 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 30 and 50) is replaced with the substituted amino acids as described above (either conservative or nonconservative).

L, S, T, M, V, N, Q, F, W, Y, P, or C; L12 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T13 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S14 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; C15 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; L16 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K17 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K18 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; R19 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E20 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E21 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; M22 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K23 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L24 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K25 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E26 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; C27 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; V28 replaced with D, E, H, K, R, N, O, F, W. Y, P, or C; S29 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; I30 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L31 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P32 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; R33 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K34 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E35 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; S36 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P37 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; S38 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V39 replaced with D, E, H, K, R, N, Q, F, W. Y, P, or C; R40 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; S41 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S42 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K43 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; D44 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G45 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K46 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L47 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L48 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A49 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A50 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T51 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L52 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L53 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L54 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A55 replaced with D, E, H, K, R, N,

Q, F, W, Y, P, or C; L56 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L57 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S58 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; C59 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; C60 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; L61 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T62 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V63 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V64 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S65 replaced with D, E, H, K, R, N, O, F, W, Y, P, or C, F66 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; Y67 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; Q68 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; V69 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A70 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A71 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L72 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q73 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; G74 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D75 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L76 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A77 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S78 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L79 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; R80 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A81 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E82 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L83 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q84 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; G85 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; H86 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; H87 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A88 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E89 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K90 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L91 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P92 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; A93 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G94 replaced with D, E, H, K, R, N, Q, F, W. Y, P, or C; A95 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G96 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A97 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P98 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; K99 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A100 replaced with D,

E, H, K, R, N, Q, F, W, Y, P, or C; G101 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L102 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E103 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E104 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A105 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P106 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; A107 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V108 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T109 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A110 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G111 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L112 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K113 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; I114 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F115 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; E116 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; P117 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; P118 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; A119 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P120 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; G121 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E122 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G123 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; N124 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; S125 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S126 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q127 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; N128 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; S129 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; R130 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; N131 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; K132 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; R133 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; A134 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V135 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q136 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; G137 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P138 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; E139 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E140 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T141 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G142 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C;

S143 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y144 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; T145 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F146 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; V147 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P148 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; W149 replaced with D, E, H, K, R, N, O, A, G, I, L, S, T, M, V, P, or C; L150 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L151 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S152 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F153 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; K154 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; R155 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G156 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S157 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A158 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L159 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E160 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E161 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K162 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E163 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; N164 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; K165 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; I166 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L167 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; V168 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K169 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E170 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T171 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G172 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y173 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; F174 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; F175 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; 1176 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y177 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; G178 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q179 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; V180 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L181 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y182 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; T183 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D184 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W. Y, P, or C; K185 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T186

replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Y187 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; A188 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; M189 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G190 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; H191 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L192 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; I193 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q194 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; R195 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K196 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; K197 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; V198 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; H199 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; V200 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F201 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; G202 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D203 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E204 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L205 replaced with D, E, H, K, R, N, O, F, W, Y, P, or C; S206 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C, L207 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C, V208 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T209 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L210 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F211 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; R212 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; C213 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; I214 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q215 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; N216 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; M217 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P218 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; E219 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; T220 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L221 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P222 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; N223 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; N224 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; S225 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; C226 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or P; Y227 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; S228 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C;

A229 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; G230 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; I231 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A232 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K233 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L234 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; E235 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E236 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G237 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D238 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E239 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L240 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q241 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; L242 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A243 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; I244 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; P245 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, or C; R246 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; E247 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; N248 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C; A249 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; Q250 replaced with D, E, H, K, R, A, G, I, L, S, T, M, V, F, W, Y, P, or C, 1251 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; S252 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L253 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D254 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; G255 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; D256 replaced with H, K, R, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; V257 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; T258 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; F259 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; F260 replaced with D, E, H, K, R, N, Q, A, G, I, L, S, T, M, V, P, or C; G261 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; A262 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; L263 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; K264 replaced with D, E, A, G, I, L, S, T, M, V, N, Q, F, W, Y, P, or C; L265 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C; and/or L266 replaced with D, E, H, K, R, N, Q, F, W, Y, P, or C. In another embodiment, site directed changes at the amino acid level of [0186]BLyS can be made by replacing a particular amino acid with a non-conservative substitution. Antibodies of the present invention may bind BLyS amino acid sequences containing non-conservative substitution mutations of the polypeptide of any one of SEQ

ID NOS:3230-3237.

[0187] In an additional embodiment, antibodies of the present invention bind BLyS polypeptides comprising, or alternatively consisting of, a BLyS amino acid sequence in which more than one amino acid (e.g., 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 30 and 50) is replaced with the substituted amino acids as described above (either conservative or nonconservative).

[0188] Replacement of amino acids can also change the selectivity of the binding of a ligand to cell surface receptors. For example, Ostade *et al.*, *Nature 361*:266-268 (1993) describes certain mutations resulting in selective binding of TNF-alpha to only one of the two known types of TNF receptors. Since BLyS is a member of the TNF polypeptide family, mutations similar to those in TNF-alpha are likely to have similar effects in BLyS polypeptides.

[0189] Sites that are critical for ligand-receptor binding can also be determined by structural analysis such as crystallization, nuclear magnetic resonance or photoaffinity labeling (Smith et al., J. Mol. Biol. 224:899-904 (1992) and de Vos et al. Science 255:306-312 (1992)).

be made in sequences encoding amino acids in the TNF conserved domain, e.g., in positions Gly-191 through Leu-284 of SEQ ID NO:3228 or in positions Gly-172 through Leu-265 of SEQ ID NO:3229, may modulate rather than completely eliminate functional activities (e.g., biological activities) of BLyS polypeptides or fragments or variants thereof. Accordingly, antibodies of the present invention may bind BLyS polypeptides that have mutations in the TNF conserved domain. In preferred embodiments, antibodies of the present invention may bind BLyS polypeptides that have mutations in the TNF conserved domain and act as antagonists of BLyS. In other preferred embodiments, antibodies of the present invention may bind BLyS polypeptides that have mutations in the TNF conserved domain and act as agonists of BLyS.

[0191] Recombinant DNA technology known to those skilled in the art (see, for instance, DNA shuffling *supra*) can be used to create novel mutant proteins or muteins including single or multiple amino acid substitutions, deletions, additions or fusion proteins. Such modified polypeptides can show, e.g., enhanced activity or increased stability. In addition, they may be purified in higher yields and show better solubility than

the corresponding natural polypeptide, at least under certain purification and storage conditions.

Thus, the invention also encompasses antibodies that bind BLyS [0192] derivatives and analogs that have one or more amino acid residues deleted, added, or substituted to generate BLyS polypeptides, e.g., that are better suited for expression, scale up, etc., in the host cells. For example, cysteine residues can be deleted or substituted with another amino acid residue in order to eliminate disulfide bridges; N-linked glycosylation sites can be altered or eliminated to achieve, for example, expression of a homogeneous product that is more easily recovered and purified from yeast hosts which are known to hyperglycosylate N-linked sites. To this end, a variety of amino acid substitutions at one or both of the first or third amino acid positions on any one or more of the glycosylation recognition sequences in the BLyS polypeptides of the invention, and/or an amino acid deletion at the second position of any one or more such recognition sequences will prevent glycosylation of the BLyS at the modified tripeptide sequence (see, e.g., Miyajimo et al., EMBO J 5(6):1193-1197). By way of non-limiting example, mutation of the serine at position 244 to alanine either singly or in combination with mutation of the asparagine at position 242 to glutamine abolishes glycosylation of the mature soluble form of BLyS (e.g., amino acids 134-285 of SEQ ID NO:3228) when expressed in the yeast Pichea pastoris. A mutant BLyS polypeptide in which only the asparagine at position 242 is mutated to glutamine, is still gycosylated when expressed in Pichea pastoris. In this mutant, the glycosylation event may be due to the activation or unmasking of an O-linked glyscosylation site at serine 244. Similar mutations affecting glycosylation could also be made in the BLyS polypeptide of SEQ ID NO:3229, i.e., aspargine-223 to glutamine and/or serine-224 to alanine of SEQ ID NO:3229. Additionally, one or more of the amino acid residues of the polypeptides of the invention (e.g., arginine and lysine residues) may be deleted or substituted with another residue to eliminate undesired processing by proteases such as, for example, furins or kexins. One possible result of such a mutation is that BLyS polypeptide of the invention is not cleaved and released from the cell surface. Accordingly, antibodies of the invention may bind BLyS derivatives and analogs that have one or more amino acid residues deleted, added, or substituted. In other embodiments, antibodies of the invention may bind BLyS derivatives, variants or analogs that are unable to be cleaved from the cell surface.

[0193] In a specific embodiment, antibodies of the invention bind BLyS polypeptides in which Lys-132 and/or Arg-133 of the BLyS sequence shown in SEQ ID NO:3228 is mutated to another amino acid residue, or deleted altogether, to prevent or diminish release of the soluble form of BLyS from cells expressing BLyS. In a more specific embodiment, antibodies of the invention bind BLyS polypeptides in which Lys-132 of the BLyS sequence shown in SEQ ID NO:3228 is mutated to Ala-132. In another, nonexclusive specific embodiment, antibodies of the invention bind BLyS polypeptides in which Arg-133 of the BLyS sequence shown in SEQ ID NO:3228 is mutated to Ala-133. These mutated proteins, and/or have uses such as, for example, in ex vivo therapy or gene therapy, to engineer cells expressing a BLyS polypeptide that is retained on the surface of the engineered cells.

[0194] In a specific embodiment, antibodies of the invention bind BLyS polypeptides in which Cys-146 of the BLyS sequence shown in SEQ ID NO:3228 is mutated to another amino acid residue, or deleted altogether, for example, to aid preventing or diminishing oligomerization of the mutant BLyS polypeptide when expressed in an expression system. In a specific embodiment, antibodies of the invention bind BLyS polypeptides in which Cys-146 is replaced with a serine amino acid residue.

[0195] In another specific embodiment, antibodies of the invention bind BLyS polypeptides in which Cys-232 of the BLyS sequence shown in SEQ ID NO:3228 is mutated to another amino acid residue, or deleted altogether, for example, to aid preventing or diminishing oligomerization of the mutant BLyS polypeptide when expressed in an expression system. In a specific embodiment, antibodies of the invention bind BLyS polypeptides in which Cys-232 is replaced with a serine amino acid residue. Polypeptides encoding these polypeptides are also encompassed by the invention.

[0196] In yet another specific embodiment, antibodies of the invention bind BLyS polypeptides in which Cys-245 of the BLyS sequence shown in SEQ ID NO:3228 is mutated to another amino acid residue, or deleted altogether, for example, to aid preventing or diminishing oligomerization of the mutant BLyS polypeptide when expressed in an expression system. In a specific embodiment, antibodies of the invention bind BLyS polypeptides in which Cys-245 is replaced with a serine amino acid residue. Polypeptides encoding these polypeptides are also encompassed by the invention.

[0197] The polypeptides of the present invention are preferably provided in an

isolated form, and preferably are substantially purified. A recombinantly produced version of the BLyS polypeptides can be substantially purified by the one-step method described in Smith and Johnson, *Gene* 67:31-40 (1988).

The antibodies of the present invention bind BLyS polypeptides including the complete polypeptide encoded by the deposited cDNA (ATCC Deposit No. 97768) including the intracellular, transmembrane and extracellular domains of the polypeptide encoded by the deposited cDNA, the mature soluble polypeptide encoded by the deposited cDNA, the extracellular domain minus the intracellular and transmembrane domains of the protein, the complete polypeptide of SEQ ID NO:3228, the mature soluble polypeptide of SEQ ID NO:3228, e.g., amino acids 134-285 of SEQ ID NO:3228, the extracellular domain of SEQ ID NO:3228, amino acid residues 73-285 of SEQ ID NO:3228 minus the intracellular and transmembrane domains, as well as polypeptides which have at least 80%, 85%, 90% similarity, more preferably at least 95% similarity, and still more preferably at least 96%, 97%, 98% or 99% similarity to those described above. Polynucleotides encoding these polypeptides are also encompassed by the invention.

[0199] The antibodies of the present invention bind BLyS polypeptides including the complete polypeptide encoded by the deposited cDNA including the intracellular, transmembrane and extracellular domains of the polypeptide encoded by the deposited cDNA (ATCC Deposit No. 203518), the mature soluble polypeptide encoded by the deposited cDNA, the extracellular domain minus the intracellular and transmembrane domains of the protein, the complete polypeptide of SEQ ID NO:3229, the mature soluble of SEQ ID NO:3229, e.g., amino acid residues 134-266 of SEQ ID NO:3229, the extracellular domain of SEQ ID NO:3229, e.g., amino acid residues 73-266 of SEQ ID NO:3229 minus the intracellular and transmembrane domains, as well as polypeptides which have at least 80%, 85%, 90% similarity, more preferably at least 95% similarity, and still more preferably at least 96%, 97%, 98% or 99% similarity to those described above. Polynucleotides encoding these polypeptides are also encompassed by the invention.

[0200] Further antibodies of the present invention bind polypeptides including polypeptides at least 80%, or at least 85% identical, more preferably at least 90% or 95% identical, still more preferably at least 96%, 97%, 98% or 99% identical to the polypeptide encoded by the deposited cDNA (ATCC Deposit No. 97768) or to the polypeptide of SEQ

ID NO:3228, and also include antibodies that bind portions of such polypeptides with at least 30 amino acids and more preferably at least 50 amino acids.

[0201] Further antibodies of the present invention bind polypeptides including polypeptides at least 80%, or at least 85% identical, more preferably at least 90% or 95% identical, still more preferably at least 96%, 97%, 98% or 99% identical to the polypeptide encoded by the deposited cDNA (ATCC Deposit No. 203518) or to the polypeptide of SEQ ID NO:3229, and also include antibodies that bind portions of such polypeptides with at least 30 amino acids and more preferably at least 50 amino acids. Polynucleotides encoding these polypeptides are also encompassed by the invention.

[0202] By "% similarity" for two polypeptides is intended a similarity score produced by comparing the amino acid sequences of the two polypeptides using the Bestfit program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711) and the default settings for determining similarity. Bestfit uses the local homology algorithm of Smith and Waterman (Advances in Applied Mathematics 2:482-489, 1981) to find the best segment of similarity between two sequences.

[0203] By a polypeptide having an amino acid sequence at least, for example, 95% "identical" to a reference amino acid sequence of a BLyS polypeptide is intended that the amino acid sequence of the polypeptide is identical to the reference sequence except that the polypeptide sequence may include up to five amino acid alterations per each 100 amino acids of the reference amino acid of the BLyS polypeptide. In other words, to obtain a polypeptide having an amino acid sequence at least 95% identical to a reference amino acid sequence, up to 5% of the amino acid residues in the reference sequence may be deleted or substituted with another amino acid, or a number of amino acids up to 5% of the total amino acid residues in the reference sequence may be inserted into the reference sequence. These alterations of the reference sequence may occur at the amino or carboxy terminal positions of the reference amino acid sequence or anywhere between those terminal positions, interspersed either individually among residues in the reference sequence or in one or more contiguous groups within the reference sequence.

[0204] As a practical matter, whether any particular polypeptide is at least 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% identical to, for instance, the amino acid sequence of SEQ ID NO:3228, the amino acid sequence encoded by the deposited cDNA

clone HNEDU15 (ATCC Accession No. 97768), or fragments thereof, or, for instance, to the amino acid sequence of SEQ ID NO:3229, the amino acid sequence encoded by the deposited cDNA clone HDPMC52 (ATCC Accession No. 203518), or fragments thereof, can be determined conventionally using known computer programs such the Bestfit program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711). When using Bestfit or any other sequence alignment program to determine whether a particular sequence is, for instance, 95% identical to a reference sequence according to the present invention, the parameters are set, of course, such that the percentage of identity is calculated over the full length of the reference amino acid sequence and that gaps in homology of up to 5% of the total number of amino acid residues in the reference sequence are allowed.

In a specific embodiment, the identity between a reference (query) [0205] sequence (a sequence of the present invention) and a subject sequence, also referred to as a global sequence alignment, is determined using the FASTDB computer program based on the algorithm of Brutlag et al. (Comp. App. Biosci. 6:237-245 (1990)). Preferred parameters used in a FASTDB amino acid alignment are: Matrix=PAM 0, k-tuple=2, Mismatch Penalty=1, Joining Penalty=20, Randomization Group Length=0, Cutoff Score=1, Window Size=sequence length, Gap Penalty=5, Gap Size Penalty=0.05, Window Size=500 or the length of the subject amino acid sequence, whichever is shorter. According to this embodiment, if the subject sequence is shorter than the query sequence due to N- or C-terminal deletions, not because of internal deletions, a manual correction is made to the results to take into consideration the fact that the FASTDB program does not account for N- and C-terminal truncations of the subject sequence when calculating global percent identity. For subject sequences truncated at the N- and C-termini, relative to the query sequence, the percent identity is corrected by calculating the number of residues of the query sequence that are N- and C-terminal of the subject sequence, which are not matched/aligned with a corresponding subject residue, as a percent of the total bases of the query sequence. A determination of whether a residue is matched/aligned is determined by results of the FASTDB sequence alignment. This percentage is then subtracted from the percent identity, calculated by the above FASTDB program using the specified parameters, to arrive at a final percent identity score. This final percent identity score is

what is used for the purposes of this embodiment. Only residues to the N- and C-termini of the subject sequence, which are not matched/aligned with the query sequence, are considered for the purposes of manually adjusting the percent identity score. That is, only query residue positions outside the farthest N- and C-terminal residues of the subject sequence. For example, a 90 amino acid residue subject sequence is aligned with a 100 residue query sequence to determine percent identity. The deletion occurs at the N-terminus of the subject sequence and therefore, the FASTDB alignment does not show a matching/alignment of the first 10 residues at the N-terminus. The 10 unpaired residues represent 10% of the sequence (number of residues at the N- and C-termini not matched/total number of residues in the query sequence) so 10% is subtracted from the percent identity score calculated by the FASTDB program. If the remaining 90 residues were perfectly matched the final percent identity would be 90%. In another example, a 90 residue subject sequence is compared with a 100 residue query sequence. This time the deletions are internal deletions so there are no residues at the N- or C-termini of the subject sequence which are not matched/aligned with the query. In this case the percent identity calculated by FASTDB is not manually corrected. Once again, only residue positions outside the N- and C-terminal ends of the subject sequence, as displayed in the FASTDB alignment, which are not matched/aligned with the query sequence are manually corrected for. No other manual corrections are made for the purposes of this embodiment.

[0206] Antibodies that Immunospecifically bind BLyS Polypeptides

[0207] The present invention also encompasses antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS polypeptides, which antibodies comprise, or alternatively consist of, all or a portion of a heavy and/or light chain variable domain of the scFvs referred to in Table 1.

[0208] The present invention also encompasses methods and compositions for detecting, diagnosing and/or prognosing diseases or disorders associated with aberrant BLyS or BLyS receptor expression or inappropriate BLyS or BLyS receptor function in an animal, preferably a mammal, and most preferably a human, comprising using antibodies (including molecules which comprise, or alternatively consist of, antibody fragments or variants thereof) that immunospecifically bind to BLyS. Diseases and disorders which can

be detected, diagnosed or prognosed with the antibodies of the invention include, but are not limited to, immune disorders (e.g., lupus, rheumatoid arthritis, multiple sclerosis, myasthenia gravis, Hashimoto's disease, and immunodeficiency syndrome), inflammatory disorders (e.g., asthma, allergic disorders, and rheumatoid arthritis), infectious diseases (e.g., AIDS), and proliferative disorders (e.g., leukemia, carcinoma, and lymphoma). The present invention further encompasses methods and compositions for 102091 preventing, treating or ameliorating diseases or disorders associated with aberrant BLyS or BLyS receptor expression or inappropriate BLyS or BLyS receptor function in an animal. preferably a mammal, and most preferably a human, comprising administering to said animal an effective amount of one or more antibodies (including molecules which comprise, or alternatively consist of, antibody fragments or variants thereof) that immunospecifically bind to BLyS. Diseases and disorders which can be prevented, treated or inhibited by administering an effective amount of one or more antibodies or molecules of the invention include, but are not limited to, immune disorders (e.g., lupus, rheumatoid arthritis, multiple sclerosis, myasthenia gravis, Hashimoto's disease, and immunodeficiency syndrome), inflammatory disorders (e.g., asthma, allergic disorders, and rheumatoid arthritis), infectious diseases (e.g., AIDS), and proliferative disorders (e.g., leukemia, carcinoma, and lymphoma).

[0210] Anti-BLyS Antibodies

phage display technology. Single chain antibody molecules ("scFvs") displayed on the surface of phage particles were screened to identify those scFvs that immunospecifically bind to BLyS, including the membrane-bound form and soluble form of BLyS. The present invention encompasses the scFvs and portions thereof that were identified to immunospecifically bind to BLyS, including scFvs that immunospecifically bind to the soluble form of BLyS, scFvs that immunospecifically bind to the membrane-bound form of BLyS, and scFvs that immunospecifically bind to both the soluble form and membrane-bound form of BLyS. In particular, the present invention encompasses scFvs comprising, or alternatively consisting of, the amino acid sequence of SEQ ID NOS: 1 - 2128, as referred to in Table 1. Preferably, the scFvs of the present invention comprise, or alternatively consist of, the amino acid sequence of SEQ ID NOS:1 - 46, 321 - 329, 834 -

872, 1563 - 1595, or 1881 - 1908. The scFvs include scFvs that bind to soluble BLyS (e.g., scFvs comprising, or alternatively consisting of, an amino acid sequence of SEQ ID NOS: 1563 - 1880), scFvs that bind to the membrane-bound form of BLyS (e.g., scFvs comprising, or alternatively consisting of, an amino acid sequence of SEQ ID NOS: 1881 - 2128), and scFvs that bind to both the soluble form and the membrane-bound form of BLyS (e.g., scFvs comprising, or alternatively consisting of, an amino acid sequence of SEQ ID NOS: 1 - 1562). Molecules comprising, or alternatively consisting of, fragments or variants of these scFvs, that immunospecifically bind to BLyS are also encompassed by the invention, as are nucleic acid molecules encoding these scFvs, molecules, fragments and/or variants.

In one embodiment of the present invention, scFvs that immunospecifically [0212]bind to BLyS comprise a polypeptide having the amino acid sequence of any one of the VH domains referred to in Table 1 and/or any one of the VL domains referred to in Table 1. In preferred embodiments, scFvs of the present invention comprise the amino acid sequence of a VH domain and VL domain from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention comprise the amino acid sequence of a VH domain and VL domain from different scFvs referred to in Table 1. In another embodiment, scFvs that immunospecifically bind to BLyS, comprise a polypeptide having the amino acid sequence of any one, two, three, or more of the VH CDRs referred to in Table 1 and/or any one, two, three, or more of the VL CDRs referred to in Table 1. In preferred embodiments, scFvs of the present invention comprise the amino acid sequence of a VH CDR and VL CDR from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention comprise the amino acid sequence of a VH CDR and VL CDR from different scFvs referred to in Table 1. Molecules comprising, or alternatively consisting of, antibody fragments or variants of the scFvs referred to in Table 1 that immunospecifically bind to BLyS are also encompassed by the invention, as are nucleic acid molecules encoding these scFvs, molecules, fragments and/or variants.

[0213] (Table 1 can be found at the end of the specification just prior to the claims.)

[0214] In another embodiment of the present invention, an scFv that immunospecifically binds to a soluble form of BLyS, comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:1563 – 1880 as referred to in Table 1. In a preferred embodiment, an scFv that immunospecifically binds to a soluble form of BLyS comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:1570 – 1595. In an even more preferred embodiment, an scFv that immunospecifically binds to a soluble form of BLyS comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:1563 – 1569.

[0215] In another embodiment of the present invention, an scFv that immunospecifically binds to a membrane-bound form of BLyS comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:1881 - 2128 as referred to in Table 1. In a preferred embodiment, an scFv that immunospecifically binds to a membrane-bound form of BLyS comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:1886 - 1908. In an even more preferred embodiment, an scFv that immunospecifically binds to a membrane-bound form of BLyS comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:1881 - 1885.

[0216] In another embodiment of the present invention, an scFv that immunospecifically binds to both the soluble form and membrane-bound form of BLyS comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:1 - 1562 as referred to in Table 1. In a preferred embodiment, an scFv that immunospecifically binds to both the soluble form and membrane-bound form of BLyS comprises, or alternatively consists of, the amino acid sequence of SEQ ID NOS:834 - 872. In another preferred embodiment, an scFv that immunospecifically binds to both the soluble form and membrane-bound form of BLyS comprises, or alternatively consists of, any one of the amino acids sequences of SEQ ID NOS:1 - 46 or 321 - 329. Molecules comprising, or alternatively consisting of, fragments or variants of these scFvs, that immunospecifically bind to the soluble form of BLyS and/or the membrane-bound form of BLyS are also encompassed by the invention, as are nucleic acid molecules encoding these scFvs, molecules, fragments and/or variants.

[0217] In another embodiment of the present invention, scFvs that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of any one of the VH domains contained in SEQ ID NOS:1563 –

1880 as disclosed in Table 1 and/or any one of the VL domains contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of a VH CDR and VL CDR from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having amino acid sequence of a VH CDR and VL CDR from different scFvs referred to in Table 1. In another embodiment, scFvs that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of any one, two, three, or more of the VH CDRs SEQ ID NOS:1563 - 1880 as disclosed in Table 1 and/or any one, two, three, or more of the VL CDRs contained in contained SEQ ID NOS:1563 - 1880, as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the of the amino acid sequence of a VH domain and VL domain from different scFvs referred to in Table 1. In a preferred embodiment, scFvs that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of any one of the VH CDR3s contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1 and/or any one of the VL CDR3s contained in SEQ ID NOS: 1563 - 1880 as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of a VH CDR and VL CDR from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of a VH CDR and VL CDR from different scFvs referred to in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these scFvs, that immunospecifically bind to BLyS, preferably the soluble form of BLyS, are also encompassed by the invention, as are nucleic acid molecules encoding these scFvs, molecules, fragments and/or variants.

[0218] In another embodiment of the present invention, scFvs that

immunospecifically bind to the membrane-bound form of BLyS comprise a polypeptide having the amino acid sequence of any one of the VH domains contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1 and/or any one of the VL domains contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the soluble form of BLyS, comprise a polypeptide having the amino acid sequence of a VH CDR and VL CDR from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from different scFvs referred to in Table 1. In another embodiment, scFvs that immunospecifically bind to the membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of any one, two, three, or more of the VH CDRs contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1 and/or any one, two, three, or more of the VL CDRs contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the membrane-bound form of BLyS, comprise a polypeptide having the amino acid. sequence of a VH domain and VL domain from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from different scFvs referred to in Table 1. In a preferred embodiment, scFvs that immunospecifically bind to the membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of any one of the VH CDR3s contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1 and/or any one of the VL CDR3s contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of a VH CDR and VL CDR from different scFvs referred to in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these scFvs, that immunospecifically bind to BLyS, preferably the membrane-bound form of BLyS, are

also encompassed by the invention, as are nucleic acid molecules encoding these scFvs, molecules, fragments and/or variants.

[0219] In another embodiment of the present invention, scFvs that immunospecifically bind to the soluble form and membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of any one of the VH domains contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1 and/or any one of the VL domains contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the soluble and membrane-bound forms of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the soluble form and membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from different scFvs referred to in. Table 1. In another embodiment, scFvs that immunospecifically bind to the soluble form and membrane-bound form of BLyS comprise a polypeptide having the amino acid sequence of any one, two, three, or more of the VH CDRs contained in SEQ ID NOS:1 -1562 as disclosed in Table 1 and/or any one, two, three, or more of the VL CDRs contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the soluble form and membrane-bound form of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention that immunospecifically bind to the soluble and membrane-bound forms of BLyS, comprise a polypeptide having the amino acid sequence of a VH domain and VL domain from different scFvs referred to in Table 1. In a preferred embodiment, scFvs that immunospecifically bind to the soluble and membrane-bound forms of BLyS, comprise a polypeptide having the amino acid sequence of any one of the VH CDR3s contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1 and/or any one of the VL CDR3s contained in SEQ ID NOS:1 - 1562, as disclosed in Table 1. In preferred embodiments, scFvs of the present invention that immunospecifically bind to the soluble and membrane-bound forms of BLyS, comprise a polypeptide having the amino acid sequence of a VH CDR and VL CDR from the same scFv referred to in Table 1. In alternative embodiments, scFvs of the present invention

that immunospecifically bind to the soluble and membrane-bound forms of BLyS, comprise a polypeptide having the amino acid sequence of a VH CDR and VL CDR from different scFvs referred to in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these scFvs or molecules, that immunospecifically bind to BLyS, preferably the soluble and membrane-bound forms of BLyS, are also encompassed by the invention, as are nucleic acid molecules encoding these scFvs, molecules, fragments and/or variants.

[0220] The present invention provides antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or a polypeptide fragment of BLyS. In particular, the invention provides antibodies corresponding to the scFvs referred to in Table 1, such scFvs may routinely be "converted" to immunoglobulin molecules by inserting, for example, the nucleotide sequences encoding the VH and/or VL domains of the scFv into an expression vector containing the constant domain sequences and engineered to direct the expression of the immunoglobulin molecule, as described in more detail in Example 20, *infra*.

[0221] In one embodiment, the invention provides antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) wherein said antibodies comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one of the VH domains contained in the sequences referred to in Table 1. The present invention also provides antibodies that immunospecifically bind to a polypeptide, or polypeptide fragment of BLyS, wherein said antibodies comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one, two, three, or more of the VH CDRs contained in the sequences referred to in Table 1. Molecules comprising, or alternatively consisting of, these antibodies, or antibody fragments or variants thereof, that immunospecifically bind to BLyS or a BLyS fragment are also encompassed by the invention, as are nucleic acid molecules encoding these antibodies, molecules, fragments and/or variants.

[0222] In one embodiment of the present invention, antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind BLyS, comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VH CDR referred to in Table 1. In

particular, the invention provides antibodies that immunospecifically bind BLvS. comprising, or alternatively consisting of, a polypeptide having the amino acid sequence of a VH CDR1 contained in SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885 as disclosed in Table 1. In another embodiment, antibodies that immunospecifically bind BLyS, comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VH CDR2 contained in SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885 as disclosed in Table 1. In a preferred embodiment, antibodies that immunospecifically bind BLyS, comprise, or alternatively consist of a polypeptide having the amino acid sequence of a VH CDR3 contained in SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885 as disclosed in Table 1. In yet another embodiment, antibodies that immunospecifically bind BLyS, comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VH CDR1 contained in SEQ ID NOS:834 - 872, 1570 -1595, or 1886 – 1908 as disclosed in Table 1; a VH CDR2 contained in SEQ ID NOS: SEQ ID NOS: SEQ ID NOS:834 - 872, 1570 - 1595, or 1886 - 1908; and/or a VH CDR3 contained in SEQ ID NOS: SEQ ID NOS:834 - 872, 1570 - 1595, or 1886 - 1908 as disclosed in Table 1. Preferably, antibodies of the invention comprise, or alternatively consist of, VH CDRs that are derived from the same scFv as disclosed in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies that immunospecifically bind to BLyS are also encompassed by the invention, as are nucleic acid molecules encoding these antibodies, molecules, fragments or variants.

[0223] The present invention provides antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants) that immunospecifically bind to a polypeptide, or polypeptide fragment of BLyS. In particular, the invention provides antibodies wherein said antibodies comprise, or alternatively consist of, a VL domain having an amino acid sequence of any one of the VL domains referred to in Table 1. The present invention also provides antibodies that immunospecifically bind to a polypeptide or polypeptide fragment of BLyS, wherein said antibodies comprise, or alternatively consist of, a VL CDR having an amino acid sequence of any one, two, three, or more of the VL CDRs contained in the sequences referred to in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies that immunospecifically bind to BLyS are also encompassed by the

invention, as are nucleic acid molecules encoding these antibodies, molecules, fragments or variants.

In one embodiment of the present invention, antibodies (including [0224] molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind BLyS, comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VL CDR referred to in Table 1. In particular, the invention provides antibodies that immunospecifically bind BLyS, comprising, or alternatively consisting of, a polypeptide having the amino acid sequence of a VL CDR1 contained in SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885 as disclosed in Table 1. In another embodiment, antibodies that immunospecifically bind BLyS comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VL CDR2 contained in SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885 as disclosed in Table 1. In a preferred embodiment, antibodies comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VL CDR3 contained in SEQ ID NOS: in SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885 disclosed in Table 1. In yet another embodiment, antibodies that immnospecifically bind BLyS comprise, or alternatively consist of: a polypeptide having the amino acid sequence of a VL CDR1 contained in SEQ ID NOS:834 - 872, 1570 - 1595, or 1886 - 1908 as disclosed in Table 1; a VL CDR2 SEQ ID NOS: 834 - 872, 1570 - 1595, or 1886 - 1908 as disclosed in Table 1; and a VL CDR3 contained SEQ ID NOS:834 - 872, 1570 - 1595, or 1886 - 1908 as disclosed in Table 1. Preferably, antibodies of the invention comprise, or alternatively consist of, VL CDRs that are derived from the same scFv as disclosed in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies, that immunospecifically bind to BLyS are also encompassed by the invention, as are nucleic acid molecules encoding these antibodies, molecules, fragments or variants.

[0225] The present invention also provides antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or a polypeptide fragment of BLyS, wherein said antibodies comprise, or alternatively consist of, a VH domain of one of the scFvs referred to in Table 1 combined with a VL domain of one of the scFvs referred to in Table 1, or other VL domain. The present invention further provides antibodies (including

molecules comprise, or alternatively consist of, antibody fragments or variants thereof) that immunospecifically bind to a polypeptide or a polypeptide fragment of BLyS, wherein said antibodies comprise, or alternatively consist of, a VL domain of one of the scFvs referred to in Table 1 combined with a VH domain of one of the scFvs referred to in Table 1, or other VH domain. In a preferred embodiment, antibodies that immunospecifically bind to a polypeptide or a polypeptide fragment of BLyS, comprise, or alternatively consist of, a polypeptide having the amino acid sequence of a VH domain contained SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1 and a VL domain contained in contained SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1. In a further preferred embodiment, the antibodies of the invention comprise, or alternatively consist of, a VH and a VL domain from the same scFv as disclosed in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies, that immunospecifically bind to BLyS are also encompassed by the invention, as are nucleic acid molecules encoding these antibodies, molecules, fragments or variants.

The present invention also provides antibodies (including molecules [0226] comprising, or alternatively consisting of, antibody fragments or variants) that immunospecifically bind to a polypeptide or polypeptide fragment of BLyS, wherein said antibodies comprise, or alternatively consist of, one, two, three, or more VH CDRs and one, two, three or more VL CDRs, as referred to in Table 1. In particular, the invention provides for antibodies that immunospecifically bind to a polypeptide or polypeptide fragment of BLyS, wherein said antibodies comprise, or alternatively consist of, a VH CDR1 and a VL CDR1, a VH CDR1 and a VL CDR2, a VH CDR1 and a VL CDR3, a VH CDR2 and a VL CDR1, VH CDR2 and VL CDR2, a VH CDR2 and a VL CDR3, a VH CDR3 and a VH CDR1, a VH CDR3 and a VL CDR2, a VH CDR3 and a VL CDR3, or any combination thereof, of the VH CDRs and VL CDRs referred to in Table 1. In a preferred embodiment, one or more of these combinations are from the same scFv as disclosed in Table 1. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies, that immunospecifically bind to BLyS are also encompassed by the invention, as are nucleic acid molecules encoding these antibodies, molecules, fragments or variants.

[0227] In a preferred embodiment the invention provides antibodies wherein the VH CDRX (where X=1, 2, or 3) and VL CDRY (where Y= 1, 2, or 3) are from scFvs with the same specificity (i.e., from scFvs that bind soluble BLyS, from scFvs that bind membrane-bound BLyS, or from scFvs that bind both soluble and membrane-bound BLyS. Molecules comprising, or alternatively consisting of, fragments or variants of these antibodies, that immunospecifically bind to BLyS are also encompassed by the invention. as are nucleic acid molecules encoding these antibodies, molecules, fragments or variants. The term "antibody," as used herein, refers to immunoglobulin molecules 102281 and immunologically active portions of immunoglobulin molecules, i.e., molecules that contain an antigen binding site that immunospecifically binds an antigen. As such, the term "antibody" encompasses not only whole antibody molecules, but also antibody fragments, as well as variants (including derivatives) of antibodies and antibody fragments. Antibodies of the invention include, but are not limited to, monoclonal. multispecific, human or chimeric antibodies, single chain antibodies, single chain Fvs (scFvs), Fab fragments, F(ab')₂ fragments, Fd fragments, disulfide-linked Fvs (sdFvs), antiidiotypic (anti-Id) antibodies (including, e.g., anti-Id antibodies to antibodies of the invention), and epitope-binding fragments of any of the above. The immunoglobulin molecules of the invention can be of any type (e.g., IgG, IgE, IgM, IgD, IgA and IgY), class (e.g., IgG₁, IgG₂, IgG₃, IgG₄, IgA₁ and IgA₂) or subclass of immunoglobulin molecule. The antibodies of the present invention also include molecules comprising, or alternatively consisting of, a polypeptide having an amino acid sequence of a portion of an amino acid sequence contained SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595. or 1881 – 1908. Preferably, an antibody of the invention comprises, or alternatively consists of, a polypeptide having an amino acid sequence of a VH domain, VH CDR, VL domain, or VL CDR of any one those contained in the sequences referred to in Table 1. Antibodies of the invention also include molecules comprising, or alternatively consisting of, fragments or variants of the above antibodies that immunospecifically bind BLyS. [0229] Most preferably the antibodies of the present invention are whole antibodies or antibody fragments that immunospecifically bind human BLyS. Antibody fragments of the invention that immunospecifically bind human BLyS include, but are not limited to, Fab, Fab' and F(ab')2, Fd fragments, single-chain Fvs (scFv), single-chain

antibodies, disulfide-linked Fvs (sdFvs), fragments comprising, or alternatively consisting of, either a VL or VH domain, and epitope binding fragments of any of the above.

102301 BLyS-binding antibody fragments, including single-chain antibodies, may comprise, or alternatively consist of, the variable region(s) alone or in combination with the entirety or a portion of the following: hinge region, CH1, CH2, and CH3 domains. In a preferred embodiment, the antibodies of the invention comprise, or alternatively consist of, a polypeptide that immunospecifically binds to BLyS, said polypeptides comprise, or alternatively consist of, one, two, three, four, five, six or more CDRs referred to in Table 1, preferably a polypeptide having an amino acid sequence of a VH CDR3 and/or a VL CDR3 of contained SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 -1908 as disclosed in Table 1. Most preferably, antibodies of the invention comprise, or alternatively consist of, one, two, three, four, five, six or more CDRs from the same scFv. as referred to in Table 1. The antibodies of the invention may be from any animal origin, including birds and mammals. Preferably, the antibodies are human, murine (e.g., mouse and rat), donkey, sheep, rabbit, goat, guinea pig, camel, horse, or chicken. Most preferably, the antibodies are human antibodies. As used herein, "human" antibodies include antibodies having the amino acid sequence of a human immunoglobulin and include antibodies isolated from human immunoglobulin libraries and xenomice or other organisms that have been genetically engineered to produce human antibodies. For a detailed discussion of a few of the technologies for producing human antibodies and human monoclonal antibodies and protocols for producing such antibodies, see, e.g., PCT publications WO 98/24893; WO 92/01047; WO 96/34096; WO 96/33735; European Patent No. 0 598 877; U.S. Patent Nos. 5,413,923; 5,625,126; 5,633,425; 5,569,825; 5,661,016; 5,545,806; 5,814,318; 5,885,793; 5,916,771; and 5,939,598; and Lonberg and Huszar, Int. Rev. Immunol. 13:65-93 (1995), which are incorporated by reference herein in their entirety. Human antibodies or "humanized" chimeric monoclonal antibodies can be produced using techniques described herein or otherwise known in the art. For example, methods for producing chimeric antibodies are known in the art. See, for review the following references which are hereby incorporated in their entirety: Morrison, Science 229:1202 (1985); Oi et al., BioTechniques 4:214 (1986); Cabilly et al., U.S. Patent No. 4,816,567; Taniguchi et al., EP 171496; Morrison et al., EP 173494; Neuberger et al., WO 8601533; Robinson et al., WO 8702671; Boulianne et al., Nature 312:643 (1984);

Neuberger et al., Nature 314:268 (1985). In addition, companies such as Abgenix, Inc. (Freemont, CA) and Genpharm (San Jose, CA) can be engaged to provide human antibodies directed against a selected antigen using technology similar to that described above.

[0231] The antibodies of the present invention may be monovalent, bivalent, trivalent or multivalent. For example, monovalent scFvs can be multimerized either chemically or by association with another protein or substance. An scFv that is fused to a hexahistidine tag or a Flag tag can be multimerized using Ni-NTA agarose (Qiagen) or using anti-Flag antibodies (Stratagene, Inc.).

[0232] The antibodies of the present invention may be monospecific, bispecific, trispecific or of greater multispecificity. Multispecific antibodies may be specific for different epitopes of a BLyS polypeptide, or fragment thereof, or may be specific for both a BLyS polypeptide, or fragment thereof, and a heterologous epitope, such as a heterologous polypeptide or solid support material. See, e.g., PCT publications WO 93/17715; WO 92/08802; WO 91/00360; WO 92/05793; Tutt, et al., J. Immunol. 147:60-69 (1991); U.S. Patent Nos. 4,474,893; 4,714,681; 4,925,648; 5,573,920; 5,601,819; Kostelny et al., J. Immunol. 148:1547-1553 (1992).

[0233] The antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may bind immunospecifically to murine BLyS (e.g., a polypeptide having the amino acid sequence of human BLyS (SEQ ID NOS:3228 and/or 3229) or BLyS expressed on human monocytes; murine BLyS (SEQ ID NOS:3230 and/or 3231) or BLyS expressed on murine monocytes; rat BLyS (either the soluble forms as given in SEQ ID NOS:3232, 3233, 3234 and/or 3235 or in a membrane associated form, e.g., on the surface of rat monocytes); or monkey BLyS (e.g., the monkey BLyS polypeptides of SEQ ID NOS:3236 and/or 3237, the soluble form of monkey BLyS, or BLyS expressed on monkey monocytes), preferably the antibodies of the invention bind immunospecifically to human BLyS. Preferably, the antibodies of the invention bind immunospecifically to human and monkey BLyS. Also preferably, the antibodies of the invention bind immunospecifically to human BLyS and murine BLyS. More preferably, antibodies of the invention, bind immunospecifically and with higher affinity to human BLyS than to murine BLyS.

[0234] Antibodies of the present invention may also be described or specified in terms of their cross-reactivity. Antibodies that do not bind any other analog, ortholog, or homolog of a polypeptide of the present invention are included. Antibodies that bind polypeptides with at least 95%, at least 90%, at least 85%, at least 80%, at least 75%, at least 70%, at least 65%, at least 60%, at least 55%, and at least 50% identity (as calculated using methods known in the art and described herein) to a polypeptide of the present invention are also included in the present invention. In a specific embodiment, antibodies of the present invention cross react with APRIL (SEQ ID NO:3239; GenBank Accession No. AF046888; J. Exp. Med. 188(6):1185-1190; PCT International Publication WO97/33902). In specific embodiments, antibodies of the present invention cross-react with murine, rat and/or rabbit homologs of human proteins and the corresponding epitopes thereof. Antibodies that do not bind polypeptides with less than 95%, less than 90%, less than 85%, less than 80%, less than 75%, less than 70%, less than 65%, less than 60%, less than 55%, and less than 50% identity (as calculated using methods known in the art and described herein) to a polypeptide of the present invention are also included in the present invention. In a specific embodiment, the above-described cross-reactivity is with respect to any single specific antigenic or immunogenic polypeptide, or combination(s) of 2, 3, 4, 5, or more of the specific antigenic and/or immunogenic polypeptides disclosed herein. Further included in the present invention are antibodies which bind polypeptides encoded by polynucleotides which hybridize to a polynucleotide of the present invention under hybridization conditions (as described herein).

[0235] In preferred embodiments, the antibodies of the present invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), immunospecifically bind to BLyS and do not cross-react with any other antigens. In more preferred embodiments, the antibodies of the invention immunopecifically bind to BLyS and do not cross-react with TRAIL, APRIL, Endokine-alpha, TNF-alpha, TNF-beta, Fas-L or LIGHT.

[0236] The present invention also provides for a nucleic acid molecule, generally

isolated, encoding an antibody of the invention (including molecules [0237] comprising, or alternatively consisting of, antibody fragments or variants thereof). In one embodiment, a nucleic acid molecule of the invention encodes an antibody comprising, or alternatively consisting of, a VH domain having an amino acid sequence of any one of the VH domains referred to in Table 1. In another embodiment, a nucleic acid molecule of the present invention encodes an antibody comprising, or alternatively consisting of, a VH CDR1 having an amino acid sequence of any one of the VH CDR1s referred to in Table 1. In another embodiment, a nucleic acid molecule of the present invention encodes an antibody comprising, or alternatively consisting of, a VH CDR2 having an amino acid sequence of any one of the VH CDR2s referred to in Table 1. In yet another embodiment, a nucleic acid molecule of the present invention encodes an antibody comprising, or alternatively consisting of, a VH CDR3 having an amino acid sequence of any one of the VH CDR3s referred to in Table 1. Nucleic acid molecules encoding antibodies that immunospecifically bind BLyS and comprise, or alternatively consist of, fragments or variants of the VH domains and/or VH CDRs are also encompassed by the invention.

In another embodiment, a nucleic acid molecule of the invention encodes [0238] an antibody (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), comprising, or alternatively consisting of, a VL domain having an amino acid sequence of any one of the VL domains referred to in Table 1. In another embodiment, a nucleic acid molecule of the present invention encodes an antibody comprising, or alternatively consisting of, a VL CDR1 having amino acid sequence of any one of the VL CDR1s referred to in Table 1. In another embodiment, a nucleic acid molecule of the present invention encodes an antibody comprising, or alternatively consisting of, a VL CDR2 having an amino acid sequence of any one of the VL CDR2s referred to in Table 1. In yet another embodiment, a nucleic acid molecule of the present invention encodes an antibody comprising, or alternatively consisting of, a VL CDR3 having an amino acid sequence of any one of the VL CDR3s referred to in Table 1. Nucleic acid encoding antibodies that immunospecifically bind BLyS and comprise, or alternatively consist of, fragments or variants of the VL domains and/or VLCDR(s) are also encompassed by the invention.

[0239] In another embodiment, a nucleic acid molecule of the invention encodes an antibody (including molecules comprising, or alternatively consisting of, antibody

fragments or variants thereof), comprising, or alternatively consisting of, a VH domain having an amino acid sequence of any one of the VH domains referred to in Table 1 and a VL domain having an amino acid sequence of any one of the VL domains referred to in Table 1. In another embodiment, a nucleic acid molecule of the invention encodes an antibody comprising, or alternatively consisting of, a VH CDR1, a VL CDR1, a VH CDR2, a VL CDR2, a VH CDR3, a VL CDR3, or any combination thereof having an amino acid sequence referred to in Table 1. Nucleic acid encoding antibodies that immunospecifically bind BLyS and comprise, or alternatively consist of, fragments or variants of the VL and/or domains and/or VHCDR(s) and/or VLCDR(s) are also encompassed by the invention.

The present invention also provides antibodies that comprise, or [0240] alternatively consist of, variants (including derivatives) of the VH domains, VH CDRs. VL domains, and VL CDRs described herein, which antibodies immunospecifically bind to BLyS. Standard techniques known to those of skill in the art can be used to introduce mutations in the nucleotide sequence encoding a molecule of the invention, including, for example, site-directed mutagenesis and PCR-mediated mutagenesis which result in amino acid substitutions. Preferably, the variants (including derivatives) encode less than 50 amino acid substitutions, less than 40 amino acid substitutions, less than 30 amino acid substitutions, less than 25 amino acid substitutions, less than 20 amino acid substitutions. less than 15 amino acid substitutions, less than 10 amino acid substitutions, less than 5 amino acid substitutions, less than 4 amino acid substitutions, less than 3 amino acid substitutions, or less than 2 amino acid substitutions relative to the reference VH domain, VHCDR1, VHCDR2, VHCDR3, VL domain, VLCDR1, VLCDR2, or VLCDR3. In specific embodiments, the variants encode substitutions of VHCDR3. In a preferred. embodiment, the variants have conservative amino acid substitutions at one or more predicted non-essential amino acid residues. A "conservative amino acid substitution" is one in which the amino acid residue is replaced with an amino acid residue having a side chain with a similar charge. Families of amino acid residues having side chains with similar charges have been defined in the art. These families include amino acids with basic side chains (e.g., lysine, arginine, histidine), acidic side chains (e.g., aspartic acid, glutamic acid), uncharged polar side chains (e.g., glycine, asparagine, glutamine, serine, threonine, tyrosine, cysteine), nonpolar side chains (e.g., alanine, valine, leucine,

isoleucine, proline, phenylalanine, methionine, tryptophan), beta-branched side chains (e.g., threonine, valine, isoleucine) and aromatic side chains (e.g., tyrosine, phenylalanine, tryptophan, histidine). Alternatively, mutations can be introduced randomly along all or part of the coding sequence, such as by saturation mutagenesis, and the resultant mutants can be screened for biological activity to identify mutants that retain activity (e.g., the ability to bind BLyS). Following mutagenesis, the encoded protein may routinely be expressed and the functional and/or biological activity of the encoded protein, (e.g., ability to immunospecifically bind BLyS) can be determined using techniques described herein or by routinely modifying techniques known in the art.

[0241] The antibodies of the invention include derivatives (i.e., variants) that are modified, e.g., by the covalent attachment of any type of molecule to the antibody such that covalent attachment does not affect the ability of the antibody to immunospecifically bind to BLyS. For example, but not by way of limitation, derivatives of the invention include antibodies that have been modified, e.g., by glycosylation, acetylation, pegylation, phosphorylation, amidation, derivatization by known protecting/blocking groups, proteolytic cleavage, linkage to a cellular ligand or other protein, etc. Any of numerous chemical modifications may be carried out by known techniques, including, but not limited to, specific chemical cleavage, acetylation, formylation, metabolic synthesis of tunicamycin, etc. Additionally, the derivative may contain one or more non-classical amino acids.

[0242] In a specific embodiment, an antibody of the invention (including a molecule comprising, or alternatively consisting of, an antibody fragment or variant thereof), that immunospecifically binds BLyS, comprises, or alternatively consists of, an amino acid sequence encoded by a nucleotide sequence that hybridizes to a nucleotide sequence that is complementary to that encoding one of the VH or VL domains referred to in Table 1 under stringent conditions, e.g., hybridization to filter-bound DNA in 6x sodium chloride/sodium citrate (SSC) at about 45° C followed by one or more washes in 0.2xSSC/0.1% SDS at about 50-65° C, under highly stringent conditions, e.g., hybridization to filter-bound nucleic acid in 6xSSC at about 45° C followed by one or more washes in 0.1xSSC/0.2% SDS at about 68° C, or under other stringent hybridization conditions which are known to those of skill in the art (see, for example, Ausubel, F.M. et al., eds., 1989, Current Protocols in Molecular Biology, Vol. I, Green Publishing

Associates, Inc. and John Wiley & Sons, Inc., New York at pages 6.3.1-6.3.6 and 2.10.3). In another embodiment, an antibody of the invention that immunospecifically binds to BLyS, comprises, or alternatively consists of, an amino acid sequence encoded by a nucleotide sequence that hybridizes to a nucleotide sequence that is complementary to that encoding one of the VH CDRs or VL CDRs referred to in Table 1 under stringent conditions, e.g., hybridization under conditions as described above, or under other stringent hybridization conditions which are known to those of skill in the art. In another embodiment, an antibody of the invention that immunospecifically binds to BLyS, comprises, or alternatively consists of, an amino acid sequence encoded by a nucleotide sequence that hybridizes to a nucleotide sequence that is complementary to that encoding one of the VH CDR3s referred to in Table 1 under stringent conditions e.g., hybridization under conditions as described above, or under other stringent hybridization conditions which are known to those of skill in the art. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

In another embodiment, an antibody (including a molecule comprising, or [0243] alternatively consisting of, an antibody fragment or variant thereof), that immunospecifically binds to BLyS comprises, or alternatively consists of, a polypeptide having an amino acid sequence that is at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 99% identical, to any one of the VH domains referred to in Table 1. In another embodiment, an antibody of the invention that immunospecifically binds to BLyS comprises, or alternatively consists of, a polypeptide having an amino acid sequence that is at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 99% identical, to any one of the VH CDRs referred to in Table 1. In another embodiment, an antibody of the invention that immunospecifically binds to BLyS comprises, or alternatively consists of, a polypeptide having an amino acid sequence that is at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 99% identical to any one of the VH CDR3s referred to in Table 1. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

[0244] In another embodiment, an antibody of the invention (including a molecule comprising, or alternatively consisting of, an antibody fragment or variant thereof), that immunospecifically binds to BLyS comprises, or alternatively consists of, a polypeptide having an amino acid sequence that is at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 99% identical, to any one of the VL domains referred to in Table 1. In another embodiment, an antibody of the invention that immunospecifically binds to BLyS comprises, or alternatively consists of, a polypeptide having an amino acid sequence that is at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 99% identical, to any one of the VL CDRs referred to in Table 1. In another embodiment, an antibody of the invention that immunospecifically binds to BLyS comprises, or alternatively consists of, a polypeptide having an amino acid sequence that is at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 99% identical, to any one of the VL CDR3s referred to in Table 1. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

alternatively consisting of, antibody fragments or variants thereof) may also be described or specified in terms of their binding affinity for to BLyS polypeptides or fragments or variants of BLyS polypeptides (e.g., to the soluble form of BLyS and/or membrane-bound form of BLyS). In specific embodiments, antibodies of the invention bind BLyS polypeptides, or fragments or variants thereof, with a dissociation constant or K_D of less than or equal to 5 X 10⁻² M, 10⁻² M, 5 X 10⁻³ M, 10⁻³ M, 5 X 10⁻⁴ M, 10⁻⁴ M, 5 X 10⁻⁵ M, or 10⁻⁵ M. More preferably, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with a dissociation constant or K_D less than or equal to 5 X 10⁻⁶ M, 10⁻⁶ M, 5 X 10⁻⁷ M, 10⁻⁷ M, 5 X 10⁻⁸ M, or 10⁻⁸ M. Even more preferably, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with a dissociation constant or K_D less than or equal to 5 X 10⁻⁹ M, 10⁻⁹ M, 5 X 10⁻¹⁰ M, 10⁻¹⁰ M, 5 X 10⁻¹¹ M, 10⁻¹¹ M, 5 X 10⁻¹² M, 10⁻¹² M, 5 X -13 M, 10⁻¹³ M, 5 X 10⁻¹⁴ M, 10⁻¹⁴ M, 5 X 10⁻¹⁵ M. The invention encompasses antibodies that bind BLyS polypeptides

with a dissociation constant or K_D that is within any one of the ranges that are between each of the individual recited values.

[0246] In specific embodiments, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with an off rate (k_{off}) of less than or equal to 5 X 10⁻² sec⁻¹, 10⁻² sec⁻¹, 5 X 10⁻³ sec⁻¹ or 10⁻³ sec⁻¹. More preferably, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with an off rate (k_{off}) less than or equal to 5 X 10⁻⁴ sec⁻¹, 10⁻⁴ sec⁻¹, 5 X 10⁻⁵ sec⁻¹, or 10⁻⁵ sec⁻¹5 X 10⁻⁶ sec⁻¹, 10⁻⁶ sec⁻¹, 5 X 10⁻⁷ sec⁻¹ or 10⁻⁷ sec⁻¹. The invention encompasses antibodies that bind BLyS polypeptides with an off rate (k_{off}) that is within any one of the ranges that are between each of the individual recited values.

In other embodiments, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with an on rate (k_{on}) of greater than or equal to 10^3 M⁻¹ sec⁻¹, 5×10^3 M⁻¹ sec⁻¹, 10^4 M⁻¹ sec⁻¹ or 5×10^4 M⁻¹ sec⁻¹. More preferably, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with an on rate (k_{on}) greater than or equal to 10^5 M⁻¹ sec⁻¹, 5×10^5 M⁻¹ sec⁻¹, 10^6 M⁻¹ sec⁻¹, or 5×10^6 M⁻¹ sec⁻¹ or 10^7 M⁻¹ sec⁻¹. The invention encompasses antibodies that bind BLyS polypeptides with on rate (k_{on}) that is within any one of the ranges that are between each of the individual recited values.

[0248] The invention also encompasses antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that have one or more of the same biological characteristics as one or more of the antibodies described herein. By "biological characteristics" is meant, the in vitro or in vivo activities or properties of the antibodies, such as, for example, the ability to bind to BLyS (e.g., the soluble form of BLyS, the membrane-bound form of BLyS, the soluble form and membrane-bound form of BLyS), and/or an antigenic and/or epitope region of BLyS), the ability to substantially block BLyS/BLyS receptor (e.g., TACI - GenBank accession number AAC51790 and/or BCMA -GenBank accession number NP_001183) binding, or the ability to block BLyS mediated biological activity (e.g., stimulation of B cell proliferation and immunoglobulin production). Optionally, the antibodies of the invention will bind to the same epitope as at least one of the antibodies specifically referred to herein. Such epitope binding can be routinely determined using assays known in the art.

The present invention also provides for antibodies (including molecules [0249] comprising, or alternatively consisting of, antibody fragments or variants thereof), that neutralize BLyS or a fragment thereof, said antibodies comprising, or alternatively consisting of, a portion (i.e., a VH domain, VL domain, VH CDR1, VH CDR2, VH CDR3, VL CDR1, VL CDR2, or VL CDR3) of an scFv referred to in Table 1, more preferably having an amino acid sequence contained in SEQ ID NOS:834 - 872, 1570 -1595, or 1886 - 1908, and even more preferably having an amino acid sequence contained in SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885 as disclosed in Table 1, or a fragment or variant thereof. By an antibody that "neutralizes BLyS or a fragment thereof' is meant an antibody that diminishes or abolishes the ability of BLyS to bind to its receptor (e.g., TACI and BCMA) to stimulate B cell proliferation, to stimulate immunoglobulin secretion by B cells, and/or to stimulate the BLyS receptor signalling cascade. In one embodiment, an antibody that neutralizes BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that neutralizes BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that neutralizes BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH CDR domain in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In a preferred embodiment, an antibody that neutralizes BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH CDR3 contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that neutralizes BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL CDR domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 -1908 as disclosed in Table 1, or a fragment or variant thereof. In another preferred embodiment, an antibody that neutralizes BLyS or a fragment thereof, comprises, or

alternatively consists of, a polypeptide having the amino acid sequence of a VL CDR3 contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

The present invention also provides for antibodies (including molecules [0250] comprising, or alternatively consisting of, antibody fragments or variants thereof), that inhibit (i.e., diminish or abolish) BLyS mediated B cell proliferation as determined by any method known in the art such as, for example, the assays described in Examples 21 and 22, infra, said antibodies comprising, or alternatively consisting of, a portion (e.g., a VH domain, VL domain, VH CDR1, VH CDR2, VH CDR3, VL CDR1, VL CDR2, or VL CDR3) of an scFv having an amino acid sequence SEQ ID NOS:834 - 872, 1570 - 1595, 1886 - 1908, and even more preferably having an amino acid sequence SEQ ID NOS:1 -46, 321 - 329, 1563 - 1569, 1881 - 1885 as disclosed in Table 1 or a fragment or variant thereof. In one embodiment, an antibody that inhibits BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 -1595, or 1881 - 1908, as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that inhibits BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In a preferred embodiment, an antibody that inhibits BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH CDR3 contained in SEO ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another preferred embodiment, an antibody that inhibits BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL CDR3 contained SEQ ID NOS:1 -46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

[0251] The present invention also provides for antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), that

enhance the activity of BLyS or a fragment thereof, said antibodies comprising, or alternatively consisting of, a portion (i.e., a VH domain, VL domain, VH CDR1, VH CDR2, VH CDR3, VL CDR1, VL CDR2, or VL CDR3) of an scFv having an amino acid sequence SEQ ID NOS:834 - 872, 1570 - 1595, or 1886 - 1908, and preferably having an amino acid sequence of SEQ ID NOS:1 - 46, 321 - 329, 1563 - 1569, or 1881 - 1885, as disclosed in Table 1, or a fragment or variant thereof. By an antibody that "enhances the activity of BLyS or a fragment thereof" is meant an antibody increases the ability of BLyS to bind to its receptor (e.g., TACI or BCMA), to stimulate B cell proliferation, to stimulate immunoglobulin secretion by B cells, and/or to stimulate the BLyS receptor signalling cascade. In one embodiment, an antibody that enhances the activity of BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that enhances the activity of BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 -1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that enhances the activity of BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH CDR domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In a preferred embodiment, an antibody that enhances the activity of BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH CDR3 contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 -1908 as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that enhances BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL CDR domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another preferred embodiment, an antibody that enhances the activity of BLyS or a fragment thereof, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL CDR3 contained in SEQ ID NOS:1 -46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a

fragment or variant thereof. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

[0252] The present invention also provides for antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), that stimulate BLyS mediated B cell proliferation as determined by any method known in the art, such as, for example, the assays described in Examples 21 and 22, infra, said antibodies comprising, or alternatively consisting of, a portion (e.g., a VH domain, VL domain, VH CDR1, VH CDR2, VH CDR3, VL CDR1, VL CDR2, or VL CDR3) of an scFv having an amino acid sequence of SEQ ID NOS:834 - 872, 1570 - 1595, or 1886 -1908, and even more preferably having an amino acid sequence of SEO ID NOS:1 - 46. 321 - 329, 1563 - 1569, or 1881 - 1885 as disclosed in Table 1 or a fragment or variant thereof. In one embodiment, an antibody that stimulates BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 -1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another embodiment, an antibody that stimulates BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL domain contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In a preferred embodiment, an antibody that stimulates BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VH CDR3 contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. In another preferred embodiment, an antibody that stimulates BLyS mediated B cell proliferation, comprises, or alternatively consists of, a polypeptide having the amino acid sequence of a VL CDR3 contained in SEQ ID NOS:1 - 46, 321 - 329, 834 - 872, 1563 - 1595, or 1881 - 1908 as disclosed in Table 1, or a fragment or variant thereof. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

[0253] The present invention also provides for fusion proteins comprising, or alternatively consisting of, an antibody (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), that immunospecifically binds to BLyS, and a heterologous polypeptide. Preferably, the heterologous polypeptide to which

the antibody is fused to is useful for B-cell function or is useful to target the antibody to Bcells. In an alternative preferred embodiment, the heterologous polypeptide to which the antibody is fused to is useful for monocyte cell function or is useful to target the antibody to a monocyte. In another embodiment, the heterologous polypertide to which the antibody is fused is albumin (including but not limited to recombinant human serum albumin or fragments or variants thereof (see, e.g., U.S. Patent No. 5.876,969, issued March 2, 1999, EP Patent 0 413 622, and U.S. Patent No. 5,766,883, issued June 16, 1998, herein incorporated by reference in their entirety)). In a preferred embodiment, antibodies of the present invention (including fragments or variants thereof) are fused with the mature form of human serum albumin (i.e., amino acids 1 - 585 of human serum albumin as shown in Figures 1 and 2 of EP Patent 0 322 094) which is herein incorporated by reference in its entirety. In another preferred embodiment, antibodies of the present invention (including fragments or variants thereof) are fused with polypeptide fragments comprising, or alternatively consisting of, amino acid residues 1-x of human serum albumin, where x is an integer from 1 to 585 and the albumin fragment has human serum albumin activity. In another preferred embodiment, antibodies of the present invention (including fragments or variants thereof) are fused with polypeptide fragments comprising, or alternatively consisting of, amino acid residues 1-z of human serum albumin, where z is an integer from 369 to 419, as described in U.S. Patent 5,766,883 herein incorporated by reference in its entirety. Antibodies of the present invention (including fragments or variants thereof) may be fused to either the N- or C-terminal end of the heterologous protein (e.g., immunoglobulin Fc polypeptide or human serum albumin polypeptide).

[0254] In one embodiment, a fusion protein of the invention comprises, or alternatively consists of, a polypeptide having the amino acid sequence of any one or more of the VH domains referred to in Table 1 or the amino acid sequence of any one or more of the VL domains referred to in Table 1 or fragments or variants thereof, and a heterologous polypeptide sequence. In another embodiment, a fusion protein of the present invention comprises, or alternatively consists of, a polypeptide having the amino acid sequence of any one, two, three, or more of the VH CDRs referred to in Table 1, or the amino acid sequence of any one, two, three, or more of the VL CDRs referred to in Table 1, or fragments or variants thereof, and a heterologous polypeptide sequence. In a

preferred embodiment, the fusion protein comprises, or alternatively consists of, a polypeptide having the amino acid sequence of, a VH CDR3 referred to in Table 1, or fragment or variant thereof, and a heterologous polypeptide sequence, which fusion protein immunospecifically binds to BLyS. In another embodiment, a fusion protein comprises, or alternatively consists of a polypeptide having the amino acid sequence of at least one VH domain referred to in Table 1 and the amino acid sequence of at least one VL domain referred to in Table 1 or fragments or variants thereof, and a heterologous polypeptide sequence. Preferably, the VH and VL domains of the fusion protein correspond to the same scFv referred to in Table 1. In yet another embodiment, a fusion protein of the invention comprises, or alternatively consists of a polypeptide having the amino acid sequence of any one, two, three or more of the VH CDRs referred to in Table 1 and the amino acid sequence of any one, two, three or more of the VL CDRs referred to in Table 1, or fragments or variants thereof, and a heterologous polypeptide sequence. Preferably, two, three, four, five, six, or more of the VHCDR(s) or VLCDR(s) correspond to the same scFv referred to in Table 1. Nucleic acid molecules encoding these fusion proteins are also encompassed by the invention.

[0255] The present invention also provides: antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), that immunospecifically bind to the soluble form of BLyS; antibodies that immunospecifically bind to the membrane-bound form of BLyS; and antibodies that immunospecifically bind to both the soluble form and membrane-bound form of BLyS.

[0256] In one embodiment of the present invention, antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to the soluble form of BLyS, comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one or more of the VH domains contained in SEQ ID NOS:1563 – 1880 as disclosed in Table 1 and/or the amino acid sequence of any one or more of the VL domains contained in SEQ ID NOS: 1563 – 1880 as disclosed in Table 1, or fragment(s) or variant(s) (including derivative) thereof. Preferably, the VH and VL domains of the antibody correspond to the same scFv as disclosed in Table 1. In another embodiment, antibodies that immunospecifically bind to the soluble form of BLyS are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one, two,

three, or more of the VH CDRs contained SEQ ID NOS: 1563 - 1880 as disclosed in Table 1 and/or the amino acid sequence of any one, two, three, or more of the VL CDRs contained in SEQ ID NOS: 1563 - 1880 as disclosed in Table 1, or fragment(s) or variant(s) thereof. Preferably, two, three, four, five, six or more of the VH and VL CDRs of the antibody correspond to the same scFv as disclosed in Table 1. In a preferred embodiment, antibodies that immunospecifically bind to the soluble form of BLyS are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one or more of the VH CDR3s contained in SEQ ID NOS: 1563 - 1880 as disclosed in Table 1 and/or the amino acid sequence of any one or more of the VL CDR3s contained in SEQ ID NOS: 1563 - 1880 as disclosed in Table 1, or fragment(s) or variant(s) thereof. Preferably, the VHCDR3 and VLCDR3 of the antibody correspond to the same scFv, as disclosed in Table 1. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

[0257] In another embodiment of the present invention, antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to the membrane-bound form of BLyS are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one or more of the VH domains contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1 and/or the amino acid sequence of any one or more of the VL domains contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1, or a fragment or variant thereof. Preferably, the VH and VL domains of the antibody correspond to the same scFv as disclosed in Table 1. In another embodiment, antibodies that immunospecifically bind to the membrane-bound form of BLyS are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one, two, three, or more of the VH CDRs contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1 and/or the amino acid sequence of any one, two, three, or more of the VL CDRs contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1, or fragment(s) or variant(s) thereof. Preferably, two, three, four, five, six or more of the VH and VL CDRs of the antibody correspond to the same scFv as disclosed in Table 1. In a preferred embodiment, antibodies that immunospecifically bind to the membrane-bound form of BLyS are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one or more of the VH CDR3s contained in SEQ ID NOS: 1881 - 2128 as disclosed in

Table 1 and/or the amino acid sequence of any one or more of the VL CDR3s contained in SEQ ID NOS: 1881 - 2128 as disclosed in Table 1, or fragment(s) or variant(s) thereof. Preferably, the VHCDR3 and VLCDR3 of the antibody correspond to the same scFv, as disclosed in Table 1. Nucleic acid molecules encoding these antibodies are also encompassed by the invention.

[0258] In another embodiment of the present invention, antibodies (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to the soluble form and membrane-bound form of BLyS, are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one or more of the VH domains contained in SEQ ID NOS: 1 - 1562 as disclosed in Table 1 and/or the amino acid sequence of any one or more of the VL domains contained in SEQ ID NOS: 1 - 1562 as disclosed in Table 1, or a fragment or variant thereof. Preferably, the VH and VL domains of the antibody correspond to the same scFv as disclosed in Table 1. In another embodiment, antibodies that immunospecifically bind to the soluble form and membrane-bound form of BLyS are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one, two, three, or more of the VH CDRs contained in SEO ID NOS: 1 1562 as disclosed in Table 1 and/or the amino acid sequence of any one, two, three, or more of the VL CDRs contained in SEQ ID NOS: 1 - 1562 as disclosed in Table 1, or fragment(s) or variant(s) thereof. Preferably, two, three, four, five, six or more of the VH and VL CDRs of the antibody correspond to the same scFv as disclosed in Table 1. In a preferred embodiment, antibodies that immunospecifically bind to the soluble form and membrane-bound form of BLyS are provided that comprise, or alternatively consist of, a polypeptide having the amino acid sequence of any one or more of the VH CDR3s contained in SEQ ID NOS: 1 - 1562, disclosed in Table 1 and/or the amino acid sequence of any one or more of the VL CDR3s contained in SEQ ID NOS: 1 - 1562, disclosed in Table 1, or fragment(s) or variant(s) thereof. Preferably, the VHCDR3 and VLCDR3 of the antibody correspond to the same scFv, as disclosed in Table 1.

[0259] The present invention also provides for mixtures of antibodies (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS, wherein the mixture has at least one, two, three, four, five or more different antibodies of the invention. In particular,

the invention provides for mixtures of different antibodies that immunospecifically bind to the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the membrane-bound form and soluble form of BLyS. In specific embodiments, the invention provides mixtures of at least 2, preferably at least 4, at least 6, at least 8, at least 10, at least 12, at least 15, at least 20, or at least 25 different antibodies that immunospecifically bind to BLyS, wherein at least 1, at least 2, at least 4, at least 6, or at least 10, antibodies of the mixture is an antibody of the invention. In a specific embodiment, each antibody of the mixture is an antibody of the invention.

The present invention also provides for panels of antibodies (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS, wherein the panel has at least one, two, three, four, five or more different antibodies of the invention. In particular, the invention provides for panels of different antibodies that immunospecifically bind to the soluble form of BLyS, the membrane-bound form of BLyS, and/or both the membrane-bound form and soluble form of BLyS. In specific embodiments, the invention provides for panels of antibodies that have different affinities for BLyS, different specificities for BLyS, or different dissociation rates. The invention provides panels of at least 10, preferably at least 25, at least 50, at least 75, at least 100, at least 125, at least 150, at least 175, at least 200, at least 250, at least 300, at least 350, at least 400, at least 450, at least 500, at least 550, at least 600, at least 600, at least 700, at least 750, at least 800, at least 850, at least 900, at least 950, or at least 1000, antibodies. Panels of antibodies can be used, for example, in 96 well plates for assays such as ELISAs.

[0261] The present invention further provides for compositions comprising, one or more antibodies (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the invention). In one embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH domains contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR1s contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1, or

a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR2s contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1, or a variant thereof. In a preferred embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR3s contained in SEQ ID NOS:1563 - 1880, as disclosed in Table 1 or a variant thereof.

The present invention further provides for compositions comprising, one or more antibodies (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the invention). In one embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH domains contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR1s contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR2s contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1, or a variant thereof. In a preferred embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR3s contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1 or a variant thereof.

[0263] The present invention further provides for compositions comprising, one or more antibodies (including scFvs, or molecules comprising, or alternatively consisting of antibody fragments or variants of the invention). In one embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH domains contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1, or a

variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR1s contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR2s contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1, or a variant thereof. In a preferred embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH CDR3s contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1 or a variant thereof.

Other embodiments of the present invention providing for compositions 102641 comprising, one or more antibodies (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the invention) are listed below. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternative consist of, a polypeptide having an amino acid sequence of any one or more of the VL domains contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR1s contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR2s contained SEQ ID NOS:1563 - 1880 as disclosed in Table 1, or a variant thereof. In a preferred embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR3s contained in SEQ ID NOS:1563 - 1880 as disclosed in Table 1, or a variant thereof.

Other embodiments of the present invention providing for compositions [0265] comprising, one or more antibodies (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the invention) are listed below. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL domains contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR1s contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR2s SEQ ID NOS:1881 - 2128 as disclosed in Table 1, or a variant thereof. In a preferred embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR3s contained in SEQ ID NOS:1881 - 2128 as disclosed in Table 1, or a variant thereof.

[0266] Other embodiments of the present invention providing for compositions comprising, one or more antibodies (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the invention) are listed below. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL domains contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR1s contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1, or a variant thereof. In another embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid

sequence of any one or more of the VL CDR2s SEQ ID NOS:1 - 1562 as disclosed in Table 1, or a variant thereof. In a preferred embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VL CDR3s contained in SEQ ID NOS:1 - 1562 as disclosed in Table 1, or a variant thereof.

[0267] In a preferred embodiment, a composition of the present invention comprises, one, two, three, four, five, or more antibodies that comprise, or alternatively consist of, a polypeptide having an amino acid sequence of any one or more of the VH domains in disclosed in Table 1, or a variant thereof, and an amino acid sequence of any one or more of the VL domains disclosed in Table 1, or a variant thereof wherein the VH and VL domains are from scFvs with the same specificity (i.e., from scFvs that bind soluble BLyS (SEO ID NOS:1563 - 1880), from scFvs that bind membrane-bound BLyS (SEQ ID 1881 - 2128), or from scFvs that bind both soluble and membrane-bound BLyS (SEQ ID NOS:1 – 1562). In a preferred embodiment the invention provides antibodies wherein the VH CDRX (where X=1,2, or 3) and VL CDRY (where Y=1,2, or 3) are from scFvs with the same specificity (i.e., from scFvs that bind soluble BLyS (SEQ ID NOS:1563 - 1880), from scFvs that bind membrane-bound BLyS (SEQ ID NOS:1881 2128), or from scFvs that bind both soluble and membrane-bound BLyS (SEQ ID NOS:1 - 1562). In yet another embodiment, a composition of the present invention comprises one or more fusion proteins.

[0268] As discussed in more detail below, a composition of the invention may be used either alone or in combination with other compositions. The antibodies (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the present invention) may further be recombinantly fused to a heterologous polypeptide at the N- or C-terminus or chemically conjugated (including covalently and non-covalently conjugations) to polypeptides or other compositions. For example, antibodies of the present invention may be recombinantly fused or conjugated to molecules useful as labels in detection assays and effector molecules such as heterologous polypeptides, drugs, radionuclides, or toxins. See, e.g., PCT publications WO 92/08495; WO 91/14438; WO 89/12624; U.S. Patent No. 5,314,995; and EP 396,387.

[0269] Antibodies of the present invention (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the present invention) may be used, for example, but not limited to, to purify and detect BLyS, and to target the polypeptides of the present invention to cells expressing membrane-bound BLyS or BLyS receptor, including both *in vitro* and *in vivo* diagnostic and therapeutic methods. For example, the antibodies have use in immunoassays for qualitatively and quantitatively measuring levels of BLyS in biological samples. See, *e.g.*, Harlow *et al.*, Antibodies: A Laboratory Manual, (Cold Spring Harbor Laboratory Press, 2nd ed. 1988) (incorporated by reference herein in its entirety).

Methods Producing Antibodies

[0270] The antibodies of the invention (including scFvs and other molecules comprising, or alternatively consisting of antibody fragments or variants of the invention) can be produced by any method known in the art for the synthesis of antibodies, in particular, by chemical synthesis or preferably, by recombinant expression techniques.

[0271] The single chain Fvs disclosed in Table 1 were generated using phage display methods known in the art. Furthermore, other scFvs that immunospecifically bind BLyS may be generated using phage display methods known in the art. In phage display methods, functional antibody domains are displayed on the surface of phage particles which carry the polynucleotide sequences encoding them. In particular, DNA sequences encoding VH and VL domains are amplified from animal cDNA libraries (e.g., human or murine cDNA libraries of lymphoid tissues) or synthetic cDNA libraries. The DNA encoding the VH and VL domains are joined together by an scFv linker by PCR and cloned into a phagemid vector (e.g., p CANTAB 6 or pComb 3 HSS). The vector is electroporated in E. coli and the E. coli is infected with helper phage. Phage used in these methods are typically filamentous phage including fd and M13 and the VH and VL domains are usually recombinantly fused to either the phage gene III or gene VIII. Phage expressing an antigen binding domain that binds to an antigen of interest (i.e., BLyS or a fragment thereof) can be selected or identified with antigen, e.g., using labeled antigen or antigen bound or captured to a solid surface or bead. Examples of phage display methods that can be used to make the antibodies of the present invention include, but are not limited to, those disclosed in Brinkman et al., J. Immunol. Methods 182:41-50 (1995);

Ames et al., J. Immunol. Methods 184:177-186 (1995); Kettleborough et al., Eur. J. Immunol. 24:952-958 (1994); Persic et al., Gene 187 9-18 (1997); Burton et al., Advances in Immunology 57:191-280(1994); PCT application No. PCT/GB91/O1 134; PCT publications WO 90/02809; WO 91/10737; WO 92/01047; WO 92/18619; WO 93/1 1236; WO 95/15982; WO 95/20401; WO97/13844; and U.S. Patent Nos. 5,698,426; 5,223,409; 5,403,484; 5,580,717; 5,427,908; 5,750,753; 5,821,047; 5,571,698; 5,427,908; 5,516,637; 5,780,225; 5,658,727; 5,733,743 and 5,969,108; each of which is incorporated herein by reference in its entirety.

As described in the above references, after phage selection, the antibody coding regions from the phage can be isolated and used to generate whole antibodies, including human antibodies, or any other desired antigen binding fragment, and expressed in any desired host, including mammalian cells, insect cells, plant cells, yeast, and bacteria, e.g., as described below. Techniques to recombinantly produce Fab, Fab' and F(ab')2 fragments can also be employed using methods known in the art such as those disclosed in PCT publication WO 92/22324; Mullinax et al., BioTechniques 12(6):864-869 (1992); Sawai et al., AJRI 34:26-34 (1995); and Better et al., Science 240:1041-1043 (1988) (said references incorporated by reference in their entireties).

To generate whole antibodies, PCR primers including VH or VL nucleotide [0273] sequences, a restriction site, and a flanking sequence to protect the restriction site can be used to amplify the VH or VL sequences in scFv clones. Utilizing cloning techniques known to those of skill in the art, the PCR amplified VH domains can be cloned into vectors expressing a VH constant region, e.g., the human gamma 4 constant region, and the PCR amplified VL domains can be cloned into vectors expressing a VL constant region, e.g., human kappa or lambda constant regions. Preferably, the vectors for expressing the VH or VL domains comprise a promoter suitable to direct expression of the heavy and light chains in the chosen expression system, a secretion signal, a cloning site for the immunoglobulin variable domain, immunoglobulin constant domains, and a selection marker such as neomycin. The VH and VL domains may also be cloned into one vector expressing the necessary constant regions. The heavy chain conversion vectors and light chain conversion vectors are then co-transfected into cell lines to generate stable or transient cell lines that express full-length antibodies, e.g., IgG, using techniques known to those of skill in the art.

[0274] Cell lines that express antibodies that comprise the VH and VL domains of scFvs of the invention have been deposited with the American Type Culture Collection ("ATCC") on the dates listed in Table 2 and given the ATCC Deposit Numbers identified in Table 2. The ATCC is located at 10801 University Boulevard, Manassas, VA 20110-2209, USA. The ATCC deposit was made pursuant to the terms of the Budapest Treaty on the international recognition of the deposit of microorganisms for purposes of patent procedure.

Cell Line	Corresponding	SEQ ID	ATCC	ATCC Deposit
	scFv	NO:	Deposit	Date
			Number	
NSO-B11-15	I050B11-15	24	PTA-3238	March 27, 2001
NSO-anti-BLyS-6D08-18	1006D08	2	PTA-3239	March 27, 2001
NSO- anti-BLyS-116A01-60	I116A01	327	PTA-3240	March 27, 2001
IO26C04K	I026C04-K	1563	PTA-3241	March 27, 2001
IO50A12	I050A12	12	PTA-3242	March 27, 2001
IO50-B11	I050B11	9	PTA-3243	March 27, 2001

[0275] Accordingly, in one embodiment, the invention provides antibodies that comprise the VH and VL domains of scFvs of the invention.

[0276] In a preferred embodiment, an antibody of the invention is the antibody expressed by cell line NSO-B11-15.

[0277] In a preferred embodiment, an antibody of the invention is the antibody expressed by cell line NSO-anti-BLyS-6D08-18.

[0278] In a preferred embodiment, an antibody of the invention is the antibody expressed by cell line NSO- anti-BLyS-116A01-60.

[0279] In a preferred embodiment, an antibody of the invention is the antibody expressed by cell line IO26C04K.

[0280] In a preferred embodiment, an antibody of the invention is the antibody expressed by cell line IO50A12.

[0281] In a preferred embodiment, an antibody of the invention is the antibody expressed by cell line NSO-B11.

[0282] In other preferred embodiments, the invention provides antibodies that competitively inhibit binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide. In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by between 1% and 10% in a competitive inhibition assay. In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by between 1% and 10% in a competitive inhibition assay.

[0283] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 10% and up to 20% in a competitive inhibition assay.

[0284] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 20% and up to 30% in a competitive inhibition assay.

[0285] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 30% and up to 40% in a competitive inhibition assay.

[0286] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 40% and up to 50% in a competitive inhibition assay.

[0287] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 50% and up to 60% in a competitive inhibition assay.

[0288] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 60% and up to 70% in a competitive inhibition assay.

[0289] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 70% and up to 80% in a competitive inhibition assay.

[0290] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 80% and up to 90% in a competitive inhibition assay.

[0291] In preferred embodiments, the invention provides antibodies that which reduce the binding of an antibody comprising a fragment (e.g., VH domain, VL domain, VHCDR1, VHCDR2, VHCDR3, VLCDR1, VLCDR2, or VLCDR3) or variant of an scFv referred to in Table 1 to a BLyS polypeptide by at least 90% and up to 100% in a competitive inhibition assay.

[0292] In other preferred embodiments, the invention provides antibodies that competitively inhibit binding of the antibody produced by the cell line having ATCC deposit number PTA-3238 to a BLyS polypeptide.

[0293] In other preferred embodiments, the invention provides antibodies that competitively inhibit binding of the antibody produced by the cell line having ATCC deposit number PTA-3239 to a BLyS polypeptide.

[0294] In other preferred embodiments, the invention provides antibodies that competitively inhibit binding of the antibody produced by the cell line having ATCC deposit number PTA-3240 to a BLyS polypeptide.

[0295] In other preferred embodiments, the invention provides antibodies that competitively inhibit binding of the antibody produced by the cell line having ATCC deposit number PTA-3241 to a BLyS polypeptide.

[0296] In other preferred embodiments, the invention provides antibodies that competitively inhibit binding of the antibody produced by the cell line having ATCC deposit number PTA-3242 to a BLyS polypeptide.

[0297] In other preferred embodiments, the invention provides antibodies that competitively inhibit binding of the antibody produced by the cell line having ATCC deposit number PTA-3243 to a BLyS polypeptide.

For some uses, including in vivo use of antibodies in humans and in vitro [0298] detection assays, it may be preferable to use human or chimeric antibodies. Completely human antibodies are particularly desirable for therapeutic treatment of human patients. See also, U.S. Patent Nos. 4,444,887 and 4,716,111; and PCT publications WO 98/46645, WO 98/50433, WO 98/24893, WO98/16654, WO 96/34096, WO 96/33735, and WO 91/10741; each of which is incorporated herein by reference in its entirety. In a specific embodiment, antibodies of the present invention comprise one or more VH and VL domains corresponding to the human scFvs of the invention and framework regions from another immunoglobulin molecule, preferably a human immunoglobulin molecule. In a specific embodiment, antibodies of the present invention comprise one or more CDRs corresponding to the human scFvs of the invention and framework regions from another immunoglobulin molecule, preferably a human immunoglobulin molecule. In other embodiments, an antibody of the present invention comprises one, two, three, four, five, six or more VL CDRs or VH CDRs corresponding to one or more of the human scFvs referred to in Table 1, or fragments or variants thereof, and framework regions (and, optionally CDRs not derived from the scFvs in Table 1) from a human immunoglobulin

molecule. In a preferred embodiment, an antibody of the present invention comprises a VH CDR3, VL CDR3, or both, corresponding to the same scFv, or different scFvs referred to in Table 1, or fragments or variants thereof, and framework regions from a human immunoglobulin.

[0299] A chimeric antibody is a molecule in which different portions of the antibody are derived from different immunoglobulin molecules such as antibodies having a variable region derived from a human antibody and a non-human immunoglobulin constant region. Methods for producing chimeric antibodies are known in the art. See e.g., Morrison, Science 229:1202 (1985); Oi et al., BioTechniques 4:214 (1986); Gillies et al., J. Immunol, Methods 125:191-202 (1989); U.S. Patent Nos. 5,807,715; 4,816,567; and 4,816,397, which are incorporated herein by reference in their entirety. Chimeric antibodies comprising one or more CDRs from human species and framework regions from a non-human immunoglobulin molecule (e.g., framework regions from a canine or feline immunoglobulin molecule) can be produced using a variety of techniques known in the art including, for example, CDR-grafting (EP 239,400; PCT publication WO 91/09967; U.S. Patent Nos. 5,225,539; 5,530,101; and 5,585,089), veneering or resurfacing (EP 592,106; EP 519,596; Padlan, Molecular Immunology 28(4/5):489-498 (1991); Studnicka et al., Protein Engineering 7(6):805-814 (1994); Roguska et al., PNAS 91:969-973 (1994)), and chain shuffling (U.S. Patent No. 5,565,332). In a preferred embodiment, chimeric antibodies comprise a human CDR3 having an amino acid sequence of any one of the VH CDR3s or VL CDR3s referred to in Table 1, or a variant thereof, and non-human framework regions or human framework regions different from those of the frameworks in the corresponding scFv disclosed in Table 1. Often, framework residues in the framework regions will be substituted with the corresponding residue from the CDR donor antibody to alter, preferably improve, antigen binding. These framework substitutions are identified by methods well known in the art, e.g., by modeling of the interactions of the CDR and framework residues to identify framework residues important for antigen binding and sequence comparison to identify unusual framework residues at particular positions. (See, e.g., Queen et al., U.S. Patent No. 5,585,089; Riechmann et al., Nature 332:323 (1988), which are incorporated herein by reference in their entireties.)

Further, the antibodies of the invention can, in turn, be utilized to generate [0300] anti-idiotype antibodies that "mimic" BLyS polypeptides using techniques well known to those skilled in the art. (See, e.g., Greenspan & Bona, FASEB J. 7(5):437-444 (1993); and Nissinoff, J. Immunol. 147(8):2429-2438 (1991)). For example, antibodies of the invention which bind to BLyS and competitively inhibit the binding of BLyS to its receptor (as determined by assays well known in the art such as, for example, that disclosed, infra) can be used to generate antiidiotypes that "mimic" a BLvS ligand/receptor-binding domain and, as a consequence, bind to and neutralize BLyS receptors (e.g., TACI, BCMA, and TR20). Such neutralizing anti-idiotypes (including molecules comprising, or alternatively consisting of, antibody fragments or variants, such as Fab fragments of such anti-idiotypes) can be used in therapeutic regimens to neutralize BLyS. For example, such anti-idiotypic antibodies can be used to bind BLyS ligands/receptors, and thereby block BLyS mediated biological activity. Alternatively, anti-idiotypes that "mimic" a BLyS binding domain may bind to BLyS receptor(s) and induce BLyS receptor mediated signalling (e.g., activation of nuclear factor of activated T cells (NF-AT), nuclear factor-kappa B (NF-kappa B), and/or AP-1). Such agonistic antiidiotypes (including agonistic Fab fragments of these anti-idiotypes) can be used in therapeutic regimens to induce or enhance BLyS receptor mediated signalling. For example, such anti-idiotypic antibodies can be used to bind BLyS ligands/receptors, and thereby stimulate BLyS mediated biological activity (e.g., B cell proliferation and/or immunoglobulin production).

[0301] Once an antibody molecule of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) has been chemically synthesized or recombinantly expressed, it may be purified by any method known in the art for purification of an immunoglobulin molecule, or more generally, a protein molecule, such as, for example, by chromatography (e.g., ion exchange, affinity, particularly by affinity for the specific antigen after Protein A, and sizing column chromatography), centrifugation, differential solubility, or by any other standard technique for the purification of proteins. Further, the antibodies of the present invention may be fused to heterologous polypeptide sequences described herein or otherwise known in the art, to facilitate purification.

Polynucleotides Encoding an Antibody

[0302] The invention provides polynucleotides comprising, or alternatively consisting of, a nucleotide sequence encoding an antibody of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof). The invention also encompasses polynucleotides that hybridize under high stringency, or alternatively, under intermediate or lower stringency hybridization conditions, e.g., as defined supra, to polynucleotides complementary to nucleic acids having a polynucleotide sequence that encodes an antibody of the invention or a fragment or variant thereof.

[0303] The polynucleotides may be obtained, and the nucleotide sequence of the polynucleotides determined, by any method known in the art. Since the amino acid sequences of the scFv antibodies and VH domains, VL domains and CDRs thereof, are known (as described in Table 1), nucleotide sequences encoding these antibodies can be determined using methods well known in the art, i.e., the nucleotide codons known to encode the particular amino acids are assembled in such a way to generate a nucleic acid that encodes the antibody, of the invention. Such a polynucleotide encoding the antibody may be assembled from chemically synthesized oligonucleotides (e.g., as described in Kutmeier et al., BioTechniques 17:242 (1994)), which, briefly, involves the synthesis of overlapping oligonucleotides containing portions of the sequence encoding the antibody, annealing and ligating of those oligonucleotides, and then amplification of the ligated oligonucleotides by PCR.

[0304] Alternatively, a polynucleotide encoding an antibody (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may be generated from nucleic acid from a suitable source. If a clone containing a nucleic acid encoding a particular antibody is not available, but the sequence of the antibody molecule is known, a nucleic acid encoding the immunoglobulin may be chemically synthesized or obtained from a suitable source (e.g., an antibody cDNA library, or a cDNA library generated from, or nucleic acid, preferably poly A+ RNA, isolated from, any tissue or cells expressing the antibody, such as hybridoma cells selected to express an antibody of the invention) by PCR amplification using synthetic primers hybridizable to the 3' and 5' ends of the sequence or by cloning using an oligonucleotide probe specific for the particular gene sequence to identify, e.g., a cDNA clone from a cDNA library that encodes

the antibody. Amplified nucleic acids generated by PCR may then be cloned into replicable cloning vectors using any method well known in the art.

[0305] Once the nucleotide sequence of the antibody (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) is determined, the nucleotide sequence of the antibody may be manipulated using methods well known in the art for the manipulation of nucleotide sequences, e.g., recombinant DNA techniques, site directed mutagenesis, PCR, etc. (see, for example, the techniques described in Sambrook et al., 1990, Molecular Cloning, A Laboratory Manual, 2d Ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, NY and Ausubel et al., eds., 1998, Current Protocols in Molecular Biology, John Wiley & Sons, NY, which are both incorporated by reference herein in their entireties), to generate antibodies having a different amino acid sequence, for example to create amino acid substitutions, deletions, and/or insertions.

In a specific embodiment, one or more of the VH and VL domains referred [030ଗ to in Table 1, or fragments or variants thereof, is inserted within framework regions using recombinant DNA techniques known in the art. In a specific embodiment, one, two, three, four, five, six, or more of the CDRs referred to in Table 1, or fragments or variants thereof, is inserted within framework regions using recombinant DNA techniques known in the art. The framework regions may be naturally occurring or consensus framework regions, and preferably human framework regions (see, e.g., Chothia et al., J. Mol. Biol. 278: 457-479 (1998) for a listing of human framework regions, the contents of which are hereby incorporated by reference in its entirety). Preferably, the polynucleotides generated by the combination of the framework regions and CDRs encode an antibody (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that specifically binds to BLyS. Preferably, as discussed supra, polynucleotides encoding variants of antibodies or antibody fragments having one or more amino acid substitutions may be made within the framework regions, and, preferably, the amino acid substitutions improve binding of the antibody to its antigen. Additionally, such methods may be used to make amino acid substitutions or deletions of one or more variable region cysteine residues participating in an intrachain disulfide bond to generate antibody molecules, or antibody fragments or variants, lacking one or more intrachain

disulfide bonds. Other alterations to the polynucleotide are encompassed by the present invention and fall within the ordinary skill of the art.

Recombinant Expression of an Antibody

Recombinant expression of an antibody of the invention (including scFvs [0307] and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof (e.g., a heavy or light chain of an antibody of the invention or a portion thereof or a single chain antibody of the invention)), requires construction of an expression vector(s) containing a polynucleotide that encodes the antibody. Once a polynucleotide encoding an antibody molecule (e.g., a whole antibody, a heavy or light chain of an antibody, or portion thereof (preferably, but not necessarily, containing the heavy or light chain variable domain)), of the invention has been obtained, the vector(s) for the production of the antibody molecule may be produced by recombinant DNA technology using techniques well known in the art. Thus, methods for preparing a protein by expressing a polynucleotide containing an antibody encoding nucleotide sequence are described herein. Methods which are well known to those skilled in the art can be used to construct expression vectors containing antibody coding sequences and appropriate transcriptional and translational control signals. These methods include, for example, in vitro recombinant DNA techniques, synthetic techniques, and in vivo genetic recombination. The invention, thus, provides replicable vectors comprising a nucleotide sequence encoding an antibody molecule of the invention (e.g., a whole antibody, a heavy or light chain of an antibody, a heavy or light chain variable domain of an antibody, or a portion thereof, or a heavy or light chain CDR, a single chain Fv, or fragments or variants thereof), operably linked to a promoter. Such vectors may include the nucleotide sequence encoding the constant region of the antibody molecule (see, e.g., PCT Publication WO 86/05807; PCT Publication WO 89/01036; and U.S. Patent No. 5,122,464, the contents of each of which are hereby incorporated by reference in its entirety) and the variable domain of the antibody may be cloned into such a vector for expression of the entire heavy chain, the entire light chain, or both the entire heavy and light chains.

[0308] The expression vector(s) is(are) transferred to a host cell by conventional techniques and the transfected cells are then cultured by conventional techniques to produce an antibody of the invention. Thus, the invention includes host cells containing

polynucleotide(s) encoding an antibody of the invention (e.g., whole antibody, a heavy or light chain thereof, or portion thereof, or a single chain antibody of the invention, or a fragment or variant thereof), operably linked to a heterologous promoter. In preferred embodiments, for the expression of entire antibody molecules, vectors encoding both the heavy and light chains may be co-expressed in the host cell for expression of the entire immunoglobulin molecule, as detailed below.

[0309] A variety of host-expression vector systems may be utilized to express the antibody molecules of the invention. Such host-expression systems represent vehicles by which the coding sequences of interest may be produced and subsequently purified, but also represent cells which may, when transformed or transfected with the appropriate nucleotide coding sequences, express an antibody molecule of the invention in situ. These include, but are not limited to, microorganisms such as bacteria (e.g., E. coli, B. subtilis) transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing antibody coding sequences; yeast (e.g., Saccharomyces, Pichia) transformed with recombinant yeast expression vectors containing antibody coding sequences; insect cell systems infected with recombinant virus expression vectors (e.g., baculovirus) containing antibody coding sequences; plant cell systems infected with recombinant virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors (e.g., Ti plasmid) containing antibody coding sequences; or mammalian cell systems (e.g., COS, CHO, BHK, 293, 3T3 cells) harboring recombinant expression constructs containing promoters derived from the genome of mammalian cells (e.g., metallothionein promoter) or from mammalian viruses (e.g., the adenovirus late promoter; the vaccinia virus 7.5K promoter). Preferably, bacterial cells such as Escherichia coli, and more preferably, eukaryotic cells, especially for the expression of whole recombinant antibody molecule, are used for the expression of a recombinant antibody molecule. For example, mammalian cells such as Chinese hamster ovary cells (CHO), in conjunction with a vector such as the major intermediate early gene promoter element from human cytomegalovirus is an effective expression system for antibodies (Foecking et al., Gene 45:101 (1986); Cockett et al., Bio/Technology 8:2 (1990)).

[0310] In bacterial systems, a number of expression vectors may be advantageously selected depending upon the use intended for the antibody molecule being

expressed. For example, when a large quantity of such a protein is to be produced, for the generation of pharmaceutical compositions of an antibody molecule, vectors which direct the expression of high levels of fusion protein products that are readily purified may be desirable. Such vectors include, but are not limited to, the *E. coli* expression vector pUR278 (Ruther *et al.*, EMBO 1. 2:1791 (1983)), in which the antibody coding sequence may be ligated individually into the vector in frame with the lac Z coding region so that a fusion protein is produced; pIN vectors (Inouye & Inouye, Nucleic Acids Res. 13:3101-3109 (1985); Van Heeke & Schuster, J. Biol. Chem. 24:5503-5509 (1989)); and the like. pGEX vectors may also be used to express foreign polypeptides as fusion proteins with glutathione 5-transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption and binding to matrix glutathione agarose beads followed by elution in the presence of free glutathione. The pGEX vectors are designed to include thrombin or factor Xa protease cleavage sites so that the cloned target gene product can be released from the GST moiety.

[0311] In an insect system, Autographa californica nuclear polyhedrosis virus (AcNPV) may be used as a vector to express foreign genes. The virus grows in Spodoptera frugiperda cells. Antibody coding sequences may be cloned individually into non-essential regions (for example, the polyhedrin gene) of the virus and placed under control of an AcNPV promoter (for example, the polyhedrin promoter).

In mammalian host cells, a number of viral-based expression systems may be utilized. In cases where an adenovirus is used as an expression vector, the antibody coding sequence of interest may be ligated to an adenovirus transcription/translation control complex, e.g., the late promoter and tripartite leader sequence. This chimeric gene may then be inserted in the adenovirus genome by in vitro or in vivo recombination.

Insertion in a non-essential region of the viral genome (e.g., region El or E3) will result in a recombinant virus that is viable and capable of expressing the antibody molecule in infected hosts (e.g., see Logan & Shenk, Proc. Natl. Acad. Sci. USA 8 1:355-359 (1984)). Specific initiation signals may also be required for efficient translation of inserted antibody coding sequences. These signals include the ATG initiation codon and adjacent sequences. Furthermore, the initiation codon must be in phase with the reading frame of the desired coding sequence to ensure translation of the entire insert. These exogenous translational control signals and initiation codons can be of a variety of origins, both

natural and synthetic. The efficiency of expression may be enhanced by the inclusion of appropriate transcription enhancer elements, transcription terminators, etc. (see, e.g., Bittner et al., Methods in Enzymol. 153:51-544 (1987)).

[0313] In addition, a host cell strain may be chosen which modulates the expression of the inserted sequences, or modifies and processes the gene product in the specific fashion desired. Such modifications (e.g., glycosylation) and processing (e.g., cleavage) of protein products may be important for the function of the protein. Different host cells have characteristic and specific mechanisms for the post-translational processing and modification of proteins and gene products. Appropriate cell lines or host systems can be chosen to ensure the correct modification and processing of the foreign protein expressed. To this end, eukaryotic host cells which possess the cellular machinery for proper processing of the primary transcript, glycosylation, and phosphorylation of the gene product may be used. Such mammalian host cells include, but are not limited to, CHO, VERY, BHK, Hela, COS, NSO, MDCK, 293, 3T3, W138, and in particular, breast cancer cell lines such as, for example, BT483, Hs578T, HTB2, BT2O and T47D, and normal mammary gland cell line such as, for example, CRL7O3O and HsS78Bst.

[0314] For long-term, high-yield production of recombinant proteins, stable expression is preferred. For example, cell lines which stably express the antibody may be engineered. Rather than using expression vectors which contain viral origins of replication, host cells can be transformed with DNA controlled by appropriate expression control elements (e.g., promoter, enhancer, sequences, transcription terminators, polyadenylation sites, etc.), and a selectable marker. Following the introduction of the foreign DNA, engineered cells may be allowed to grow for 1-2 days in an enriched media, and then are switched to a selective media. The selectable marker in the recombinant plasmid confers resistance to the selection and allows cells to stably integrate the plasmid into their chromosomes and grow to form foci which in turn can be cloned and expanded into cell lines. This method may advantageously be used to engineer cell lines which express the antibody molecule. Such engineered cell lines may be particularly useful in screening and evaluation of compositions that interact directly or indirectly with the antibody molecule.

[0315] A number of selection systems may be used, including but not limited to, the herpes simplex virus thymidine kinase (Wigler et al., Cell 11:223 (1977)),

hypoxanthineguanine phosphoribosyltransferase (Szybalska & Szybalski, Proc. Natl. Acad. Sci. USA 48:202 (1992)), and adenine phosphoribosyltransferase (Lowy et al., Cell 22:8 17 (1980)) genes can be employed in tk-, hgprt- or aprt- cells, respectively. Also, antimetabolite resistance can be used as the basis of selection for the following genes: dhfr, which confers resistance to methotrexate (Wigler et al., Natl. Acad. Sci. USA 77:357 (1980); O'Hare et al., Proc. Natl. Acad. Sci. USA 78:1527 (1981)); gpt, which confers resistance to mycophenolic acid (Mulligan & Berg, Proc. Natl. Acad. Sci. USA 78:2072 (1981)); neo, which confers resistance to the aminoglycoside G-418 (Clinical Pharmacy 12:488-505; Wu and Wu, Biotherapy 3:87-95 (1991); Tolstoshev, Ann. Rev. Pharmacol. Toxicol. 32:573-596 (1993); Mulligan, Science 260:926-932 (1993); and Morgan and Anderson, Ann. Rev. Biochem. 62: 191-217 (1993); TIB TECH 11(5):155-2 15 (May, 1993)); and hygro, which confers resistance to hygromycin (Santerre et al., Gene 30:147 (1984)). Methods commonly known in the art of recombinant DNA technology may be routinely applied to select the desired recombinant clone, and such methods are described, for example, in Ausubel et al. (eds.), Current Protocols in Molecular Biology, John Wiley & Sons, NY (1993); Kriegler, Gene Transfer and Expression, A Laboratory Manual, Stockton Press, NY (1990); and in Chapters 12 and 13, Dracopoli et al. (eds), Current Protocols in Human Genetics, John Wiley & Sons, NY (1994); Colberre-Garapin et al., J. Mol. Biol. 150:1 (1981), which are incorporated by reference herein in their entireties. [0316] The expression levels of an antibody molecule can be increased by vector amplification (for a review, see Bebbington and Hentschel, The use of vectors based on gene amplification for the expression of cloned genes in mammalian cells in DNA cloning, Vol.3. (Academic Press, New York, 1987)). When a marker in the vector system expressing antibody is amplifiable, increase in the level of inhibitor present in culture of host cell will increase the number of copies of the marker gene. Since the amplified region is associated with the coding sequence of the antibody, production of the antibody will also increase (Crouse et al., Mol. Cell. Biol. 3:257 (1983)).

[0317] The host cell may be co-transfected with two expression vectors of the invention, the first vector encoding a heavy chain derived polypeptide and the second vector encoding a light chain derived polypeptide. The two vectors may contain identical selectable markers which enable equal expression of heavy and light chain polypeptides. Alternatively, a single vector may be used which encodes, and is capable of expressing,

both heavy and light chain polypeptides. In such situations, the light chain is preferably placed before the heavy chain to avoid an excess of toxic free heavy chain (Proudfoot, Nature 322:52 (1986); Kohler, Proc. Natl. Acad. Sci. USA 77:2 197 (1980)). The coding sequences for the heavy and light chains may comprise cDNA or genomic DNA.

[0318] Once an antibody molecule of the invention has been produced by recombinant expression, it may be purified by any method known in the art for purification of an immunoglobulin molecule, or more generally, for purification of a protein, for example, by chromatography (e.g., ion exchange, affinity, particularly by affinity for the specific antigen after Protein A, and sizing column chromatography), centrifugation, differential solubility, or by any other standard technique for the purification of proteins. Further, the antibodies of the present invention may be fused to heterologous polypeptide sequences described herein or otherwise known in the art to facilitate purification.

Antibody Characterization

[0319] Antibodies of the present invention (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may be characterized in a variety of ways. In particular, antibodies and related molecules of the invention may be assayed for the ability to immunospecifically bind to BLyS or a fragment of BLyS (e.g., to the soluble form or the membrane-bound form of BLyS) using techniques described herein or routinely modifying techniques known in the art. BLyS or BLyS fragments that may be immunospecifically bound by the compositions of the invention include, but are not limited to, human BLyS (SEQ ID NOS:3228 and/or 3229) or BLyS expressed on human monocytes; murine BLyS (SEQ ID NOS:3230 and/or 3231) or BLyS expressed on murine monocytes; rat BLyS (either the soluble forms as given in SEQ ID NOS:3232, 3233, 3234 and/or 3235 or in a membrane associated form, e.g., on the surface of rat monocytes); or monkey BLyS (e.g., the monkey BLyS polypeptides of SEQ ID NOS:3236 and/or 3237, the soluble form of monkey BLyS, or BLyS expressed on monkey monocytes) or fragments thereof. Preferably compositions of the invention bind human BLyS (SEQ ID NOS:3228 and/or 3229) or fragments thereof. Assays for the ability of the antibodies of the invention to immunospecifically bind BLyS or a fragment of BLyS may be performed in solution (e.g., Houghten, Bio/Techniques

13:412-421(1992)), on beads (e.g., Lam, Nature 354:82-84 (1991)), on chips (e.g., Fodor, Nature 364:555-556 (1993)), on bacteria (e.g., U.S. Patent No. 5,223,409), on spores (e.g., Patent Nos. 5,571,698; 5,403,484; and 5,223,409), on plasmids (e.g., Cull et al., Proc. Natl. Acad. Sci. USA 89:1865-1869 (1992)) or on phage (e.g., Scott and Smith, Science 249:386-390 (1990); Devlin, Science 249:404-406 (1990); Cwirla et al., Proc. Natl. Acad. Sci. USA 87:6378-6382 (1990); and Felici, J. Mol. Biol. 222:301-310 (1991)) (each of these references is incorporated herein in its entirety by reference). Antibodies that have been identified to immunospecifically bind to BLyS or a fragment of BLyS can then be assayed for their specificity and affinity for BLyS or a fragment of BLyS using or routinely modifying techniques described herein or otherwise known in the art.

[0320] The antibodies of the invention may be assayed for immunospecific binding to BLyS and cross-reactivity with other antigens by any method known in the art. In particular, the ability of an antibody to immunospecifically bind to the soluble form or membrane-bound form of BLyS and the specificity of the antibody, fragment, or variant for BLyS polypeptide from a particular species (e.g., murine, monkey or human, preferably human) may be determined using or routinely modifying techniques described herein or otherwise known in art.

[0321] Immunoassays which can be used to analyze immunospecific binding and cross-reactivity include, but are not limited to, competitive and non-competitive assay systems using techniques such as western blots, radioimmunoassays, ELISA (enzyme linked immunosorbent assay), "sandwich" immunoassays, immunoprecipitation assays, precipitin reactions, gel diffusion precipitin reactions, immunodiffusion assays, agglutination assays, complement-fixation assays, immunoradiometric assays, fluorescent immunoassays, and protein A immunoassays, to name but a few. Such assays are routine and well known in the art (see, e.g., Ausubel et al, eds, 1994, Current Protocols in Molecular Biology, Vol. 1, John Wiley & Sons, Inc., New York, which is incorporated by reference herein in its entirety). Exemplary immunoassays are described briefly below (but are not intended by way of limitation).

[0322] Immunoprecipitation protocols generally comprise lysing a population of cells in a lysis buffer such as RIPA buffer (1% NP-40 or Triton X-100, 1% sodium deoxycholate, 0.1% SDS, 0.15 M NaCl, 0.01 M sodium phosphate at pH 7.2, 1% Trasylol) supplemented with protein phosphatase and/or protease inhibitors (e.g., EDTA, PMSF,

aprotinin, sodium vanadate), adding the antibody of interest to the cell lysate, incubating for a period of time (e.g., 1 to 4 hours) at 40 degrees C, adding protein A and/or protein G sepharose beads to the cell lysate, incubating for about an hour or more at 40 degrees C, washing the beads in lysis buffer and resuspending the beads in SDS/sample buffer. The ability of the antibody of interest to immunoprecipitate a particular antigen can be assessed by, e.g., western blot analysis. One of skill in the art would be knowledgeable as to the parameters that can be modified to increase the binding of the antibody to an antigen and decrease the background (e.g., pre-clearing the cell lysate with sepharose beads). For further discussion regarding immunoprecipitation protocols see, e.g., Ausubel et al, eds, 1994, Current Protocols in Molecular Biology, Vol. 1, John Wiley & Sons, Inc., New York at 10.16.1.

Western blot analysis generally comprises preparing protein samples, [0323] electrophoresis of the protein samples in a polyacrylamide gel (e.g., 8%-20% SDS-PAGE depending on the molecular weight of the antigen), transferring the protein sample from the polyacrylamide gel to a membrane such as nitrocellulose, PVDF or nylon, blocking the membrane in blocking solution (e.g., PBS with 3% BSA or non-fat milk), washing the membrane in washing buffer (e.g., PBS-Tween 20), blocking the membrane with primary antibody (the antibody of interest) diluted in blocking buffer, washing the membrane in washing buffer, blocking the membrane with a secondary antibody (which recognizes the primary antibody, e.g., an anti-human antibody) conjugated to an enzymatic substrate (e.g., horseradish peroxidase or alkaline phosphatase) or radioactive molecule (e.g., ³²P or 125 I) diluted in blocking buffer, washing the membrane in wash buffer, and detecting the presence of the antigen. One of skill in the art would be knowledgeable as to the parameters that can be modified to increase the signal detected and to reduce the background noise. For further discussion regarding western blot protocols see, e.g., Ausubel et al, eds, 1994, Current Protocols in Molecular Biology, Vol. 1, John Wiley & Sons, Inc., New York at 10.8.1.

[0324] ELISAs comprise preparing antigen, coating the well of a 96-well microtiter plate with the antigen, washing away antigen that did not bind the wells, adding the antibody of interest conjugated to a detectable compound such as an enzymatic substrate (e.g., horseradish peroxidase or alkaline phosphatase) to the wells and incubating for a period of time, washing away unbound antibodies or non-specifically bound

antibodies, and detecting the presence of the antibodies specifically bound to the antigen coating the well. In ELISAs the antibody of interest does not have to be conjugated to a detectable compound; instead, a second antibody (which recognizes the antibody of interest) conjugated to a detectable compound may be added to the well. Further, instead of coating the well with the antigen, the antibody may be coated to the well. In this case, the detectable molecule could be the antigen conjugated to a detectable compound such as an enzymatic substrate (e.g., horseradish peroxidase or alkaline phosphatase). One of skill in the art would be knowledgeable as to the parameters that can be modified to increase the signal detected as well as other variations of ELISAs known in the art. For further discussion regarding ELISAs see, e.g., Ausubel et al, eds, 1994, Current Protocols in Molecular Biology, Vol. 1, John Wiley & Sons, Inc., New York at 11.2.1.

[0325] The binding affinity of an antibody (including an scFv or other molecule comprising, or alternatively consisting of, antibody fragments or variants thereof) to an antigen and the off-rate of an antibody-antigen interaction can be determined by competitive binding assays. One example of a competitive binding assay is a radioimmunoassay comprising the incubation of labeled antigen (e.g., ³H or ¹²⁵I) with the antibody of interest in the presence of increasing amounts of unlabeled antigen, and the detection of the antibody bound to the labeled antigen. The affinity of the antibody of the present invention for BLyS and the binding off-rates can be determined from the data by Scatchard plot analysis. Competition with a second antibody can also be determined using radioimmunoassays. In this case, BLyS is incubated with an antibody of the present invention conjugated to a labeled compound (e.g., ³H or ¹²⁵I) in the presence of increasing amounts of an unlabeled second anti-BLyS antibody.

[0326] In a preferred embodiment, BIAcore kinetic analysis is used to determine the binding on and off rates of antibodies (including an scFv or other molecule comprising, or alternatively consisting of, antibody fragments or variants thereof) to BLyS, or fragments of BLyS. BIAcore kinetic analysis comprises analyzing the binding and dissociation of BLyS from chips with immobilized antibodies on their surface as described in detail in Examples 6, 12, 17 and 18, *infra*.

[0327] The antibodies of the invention (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) can also be assayed for their ability to inhibit, increase, or not significantly alter, the binding of

BLyS to a BLyS receptor (e.g., TACI and BCMA) using techniques known to those of skill in the art. For example, cells expressing a receptor for BLyS (e.g., IM9, REH, ARH-77cells, Namalwa, and RPMI-8226 B cell tumor lines as wells as peripheral CD20+ B cells) can be contacted with BLyS in the presence or absence of an antibody, and the ability of the antibody to inhibit, increase, or not significantly alter, BLyS binding to the cells can be measured. BLyS binding to cells can be measured by, for example, flow cytometry or a scintillation assay. BLyS or the antibody can be labeled with a detectable compound such as a radioactive label (e.g., ³²P, ³⁵S, and ¹²⁵I) or a fluorescent label (e.g., fluorescein isothiocyanate, rhodamine, phycocythrin, phycocyanin, allophycocyanin, o-phthaldehyde and fluorescamine) to enable detection of an interaction between BLyS and a BLyS receptor and/or BLyS and an antibody of the invention. Alternatively, the ability of antibodies of the invention to inhibit, increase, or not significantly alter, BLyS binding to a BLyS receptor can be determined in cell-free assays. For example, native or recombinant BLyS (e.g., that having the amino acid sequence of amino acids 134 - 285 of SEQ ID NO:3228) or a fragment thereof can be contacted with an antibody and the ability of the antibody to inhibit, increase, or not significantly alter, BLyS from binding to a BLyS receptor can be determined. Preferably, the antibody is immobilized on a solid support and BLyS or a BLyS fragment is labeled with a detectable compound. Alternatively, BLyS or a BLyS fragment is immobilized on a solid support and the antibody is labeled with a detectable compound. BLyS may be partially or completely purified (e.g., partially or completely free of other polypeptides) or part of a cell lysate. Further, the BLyS polypeptide may be a fusion protein comprising BLyS or a biologically active portion thereof and a domain such as an Immunoglobulin Fc or glutathionine-S-transferase. For example, amino acid residues 1-154 of TACI (GenBank accession number AAC51790), or 1-48 of BCMA (GenBank accession number NP 001183) may be fused to the Fc region of an IgG molecule and used in a cell free assay to determine the ability of antibodies of the invention to inhibit, increase, or not significantly alter, BLyS binding to a BLyS receptor. Alternatively, BLyS can be biotinylated using techniques well known to those of skill in the art (e.g., biotinylation kit, Pierce Chemicals; Rockford, IL).

[0328] The antibodies of the invention (including scFvs or other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), can also

be assayed for their ability to inhibit, stimulate, or not significantly alter, BLyS-induced Bcell proliferation using techniques known to those of skill in the art. For example, B-cell proliferation can be assayed by ³H-thymidine incorporation assays and trypan blue cell counts (see, e.g., Moore et al., Science 285: 260-263 (1999)). Further, the antibodies of the invention, or fragments or variants thereof, can be assayed for their ability to block, stimulate, or not significantly alter, BLyS-induced activation of cellular signaling molecules and transcription factors such as calcium-modulator and cyclophilin ligand ("CAML"), calcineurin, nuclear factor of activated T cells transcription factor ("NF-AT"), nuclear factor-kappa B ("NF-kappa B"), and AP-1 using techniques known to those of skill in the art (see, e.g., von Bulow and Bram, Science 278:138-141(1997)). For example, NF-AT activity can be determined by electromobility gel shift assays, by detecting the expression of a protein known to be regulated by NF-AT (e.g., IL-2 expression), by detecting the induction of a reporter gene (e.g., an NF-AT regulatory element operably linked to a nucleic acid encoding a detectable marker such as luciferase, beta-galactosidase or chloramphenicol acetyltransferase (CAT)), or by detecting a cellular response (e.g., cellular differentiation, or cell proliferation).

[0329] The antibodies of the invention, or fragments or variants thereof can also be assayed for their ability to neutralize, enhance, or not significantly alter, BLyS activity. For example, antibodies or fragments or variants thereof, may be routinely tested for their ability to inhibit BLyS from binding to cells expressing the receptor for BLyS (see Example 3, infra).

Selection and Screening for Antibodies that Immunospecifically Bind to Soluble BLyS

[0330] Antibodies of the invention (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may be screened in a variety of assays to identify those antibodies that immunospecifically bind to the soluble form of BLyS. In one particular assay, antibodies that bind to the biotinylated soluble form of BLyS in solution are captured on streptavidin coated magnetic beads. This assay may be relatively applied to identify antibodies of the invention that neutralize and/or bind to BLyS. Additionally, antibodies may be assayed in neutralization assays described herein or otherwise known in the art (see Example 3, infra). For example,

antibodies may be tested for their ability to inhibit soluble BLyS (e.g., biotinylated BLyS) from binding to IM9 cells. In this assay, labeled soluble BLyS (e.g., biotinylated BLyS) is incubated with candidate anti-BLyS antibodies to allow for the formation of BLyS -anti-BLyS antibody complexes. Following incubation, an aliquot of the BLyS-anti-BLyS antibody sample is added to IM9 cells. The binding of soluble BLyS may be determined using techniques known in the art. For example, the binding of biotinylated BLyS to IM9 cells may be detected using a fluorimeter following the addition of streptavidin-delfia. Biotinylated BLyS, if it is not bound by antibodies that neutralize BLyS, binds to the cells is detected. Thus, an antibody that decreases the amount of bio-BLyS that binds to IM-9 cells (relative to a control sample in which the BLyS had been preincubated with an irrelevant antibody or no antibody at all) is identified as one that binds to and neutralizes the soluble form of BLyS. In another assay, antibodies are screened using ELISAs for those antibodies that bind to biotinylated soluble BLyS, but do not bind membrane-bound BLyS, such as, for example, BLyS on membranes from U937 cells (see Examples 2 and 9, infra). In these assays, soluble BLyS (e.g., biotinylated BLyS) and membrane-bound BLyS (e.g., on U937 membranes) are incubated in separate samples with the same antibodies and those antibodies that bind to the soluble BLyS (biotinylated BLyS), but not membrane-bound BLyS (e.g., on U937 membranes) are captured and identified. Antibodies of the invention (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may be tested to identify those antibodies that do not cross-react with APRIL, endokine-alpha, VEGI, TRAIL, TNF-alpha, TNF-beta, Fas-L, LIGHT, and PBS (see Example 4, infra). Antibodies may also be tested for their affinity for BLyS using, for example, BIAcore analysis (see Examples 6, 12, 17 and 18 infra). Antibodies may also be tested for their ability to stimulate, inhibit, or not alter, BLyS-induced immunoglobulin production and/or B-cell proliferation using techniques known to those of skill in the art. For example, human B-cells, BLyS and antibodies may be incubated together in 96 well plates and 'H-

Selection and Screening for Antibodies that Immunospecifically Bind to Membranebound BLyS

thymidine incorporation may be measured using a scintillation counter.

[0332] Antibodies of the invention (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may be screened in a variety of assays to identify those antibodies that immunospecifically bind to the membrane-bound form of BLyS. In one particular assay, antibodies that bind to BLyS on U937 membranes or immobilized histidine-tagged BLyS are captured. Other cell lines that express BLyS that might be useful for testing antibody binding to membrane-bound form of BLyS include, K-562, HL-60 and THP-1 cells. In another assay, antibodies are screened using ELISAs for those antibodies (or antibody fragments or variants) that bind to BLyS on U937 membranes or to histidine-tagged BLyS. In this assay, antibodies are added to 96 well plates coated with U937 membranes or histidine-tagged BLyS and those antibodies or antibody fragments or variants that bind to the U937 membranes or histidine-tagged BLyS are captured. In another assay, antibodies are screened using ELISAs for those antibodies (or antibody fragments or variants thereof) that do not bind to biotinylated BLyS (soluble BLyS) but bind to membrane-bound BLyS, such as, for example, that on membranes from U937 cells (see Example 2, infra). In these assays, soluble BLyS (e.g., biotinylated BLyS) and membrane-bound BLyS (e.g., on U937 membranes) are incubated in separate samples with the same antibodies (or antibody fragments or variants) and those antibodies (or antibody fragments or variants) that do not bind to the soluble BLyS (biotinylated BLyS), but bind the membrane-bound BLyS (e.g., on U937 membranes) are captured and identified. In other assays, antibodies are screened using ELISAs to determine which of the antibodies (or antibody fragments or variants) that bind to histidine-tagged BLyS or membranes from U937 cells do not cross-react with APRIL, endokine-alpha, VEGI, TRAIL, TNF-alpha, TNF-beta, Fas-L, LIGHT, and PBS (See Example 4, *infra*). ELISAs can also be used to determine which of the antibodies (or antibody fragments or variants) that bind to histidine-tagged BLyS or membranes from U937 cells bind to BLyS in the presence of TNF-alpha (see Example 4, infra). Antibodies or fragments or variants thereof that immunospecifically bind to the membrane-bound form of BLyS may also be tested for their affinity for histidine-tagged BLyS using highthroughput BIAcore analysis (see Example 14, infra).

[0333] Additionally, antibodies of the invention may be screened against cells engineered to express an "uncleavable" form of BLyS in order to determine their specificity for the membrane-bound form of BLyS. Mutations in BLyS which may

achieve this result include, but are not limited to, the mutation or deletion of amino acid residues Lys-132 and/or Arg-133 of the BLyS sequence shown in SEQ ID NO:3228. A typical mutagenesis might include mutation of one or both of residues Lys-132 or Arg-133 to alanine residues. Cells expressing such an "uncleavable" form of BLyS provide a profound reagent to use in assaying the ability of antibodies to bind the membrane-bound form of BLyS.

Selection and Screening for Antibodies that Immunospecifically Bind to Soluble and Membrane-bound BLyS

Antibodies of the invention (including scFvs and other molecules [0334] comprising, or alternately consisting of, antibody fragments or variants) may be screened in a variety of assays to identify those antibodies or antibody fragments or variants that immunospecifically bind to the soluble form and membrane-bound form of BLyS. In one particular assay, antibodies that bind to immobilized BLyS are captured. In another assay, antibodies are screened using ELISAs for those antibodies (or antibody fragments or variants) that inhibit the binding of soluble BLyS (e.g. soluble bio-BLyS) to IM-9 cells as described supra. In other assays, antibodies are screened using ELISAs for those antibodies that bind to membranes from U937 cells. Additionally, further ELISA assays may be performed using techniques known in the art to determine which antibodies do not cross-react with APRIL, endokine-alpha, VEGI, TRAIL, TNF-alpha, TNF-beta, Fas-L, LIGHT, and PBS, or those antibodies that bind to BLyS in the presence of TNF-alpha (see Example 4 infra). Antibodies may be assayed in neutralization assays using techniques described herein or otherwise known in the art. Antibodies that immunospecifically bind to the soluble and membrane-bound forms of BLyS may also be tested for their affinity for BLyS using high-throughput BIAcore analysis.

Antibody Conjugates

[0335] The present invention encompasses antibodies (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), recombinantly fused or chemically conjugated (including both covalent and non-covalent conjugations) to a heterologous polypeptide (or portion thereof, preferably at least 10, at least 20, at least 30, at least 40, at least 50, at least 60, at least 70, at least 80, at

least 90 or at least 100 amino acids of the polypeptide) to generate fusion proteins. The fusion does not necessarily need to be direct, but may occur through linker sequences. For example, antibodies of the invention may be used to target heterologous polypeptides to particular cell types (e.g., cells of monocytic lineage and B-cells), either in vitro or in vivo, by fusing or conjugating the heterologous polypeptides to antibodies of the invention that are specific for particular cell surface antigens (e.g., membrane-bound BLyS on cells of monocytic lineage) or which bind antigens that bind particular cell surface receptors (e.g., TACI and/or BCMA located on B cells). Antibodies fused or conjugated to heterologous polypeptides may also be used in in vitro immunoassays and purification methods using methods known in the art. See e.g., Harbor et al., supra, and PCT publication WO 93/2 1232; EP 439,095; Naramura et al., Immunol. Lett. 39:91-99 (1994); U.S. Patent 5,474,981; Gillies et al., PNAS 89:1428-1432 (1992); Fell et al., J. Immunol. 146:2446-2452 (1991), which are incorporated by reference in their entireties.

In one embodiment, a fusion protein comprises a polypeptide having an amino acid sequence of any one of the VH domains referred to in Table 1, and a heterologous polypeptide. In another embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VH CDR1s referred to in Table 1, and a heterologous polypeptide. In another embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VH CDR2s referred to in Table 1, and a heterologous polypeptide. In a preferred embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VH CDR3s referred to in Table 1 (i.e., SEQ ID NOS:2129 - 3227), and a heterologous polypeptide.

In another embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VL domains referred to in Table 1, and a heterologous polypeptide. In another embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VL CDR1s referred to in Table 1, and a heterologous polypeptide. In yet another embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VL CDR2s referred to in Table 1, and a heterologous polypeptide. In a preferred embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VL CDR3s referred to in Table 1, and a heterologous polypeptide.

[0338] In another embodiment, a fusion protein comprises a polypeptide having the amino acid sequence of any one of the VH domains referred to in Table 1, and one or more VL domains referred to in Table 1, and a heterologous polypeptide. In another embodiment, a fusion protein of the present invention comprises a polypeptide having the amino acid sequence of any one of the VH CDRs referred to in Table 1, and any one of the VL CDRs referred to in Table 1, and a heterologous polypeptide.

[0339] The present invention further includes compositions comprising, or alternatively consisting of, heterologous polypeptides fused or conjugated to antibody fragments. For example, the heterologous polypeptides may be fused or conjugated to a Fab fragment, Fd fragment, Fv fragment, F(ab)₂ fragment, or a portion thereof. Methods for fusing or conjugating polypeptides to antibody portions are known in the art. See, e.g., U.S. Patent Nos. 5,336,603; 5,622,929; 5,359,046; 5,349,053; 5,447,851; 5,112,946; EP 307,434; EP 367,166; PCT publications WO 96/04388; WO 9 1/06570; Ashkenazi et al., Proc. Natl. Acad. Sci. USA 88: 10535-10539 (1991); Zheng et al., J. Immunol. 154:5590-5600 (1995); and Vil et al., Proc. Natl. Acad. Sci. USA 89:11337-11341 (1992) (said references incorporated by reference in their entireties).

[0340] Additional fusion proteins of the invention may be generated through the techniques of gene-shuffling, motif-shuffling, exon-shuffling, and/or codon-shuffling (collectively referred to as "DNA shuffling"). DNA shuffling may be employed to modulate the activities of antibodies (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), such methods can be used to generate antibodies with altered activity (e.g., antibodies with higher affinities and lower dissociation rates). See, generally, U.S. Patent Nos. 5,605,793; 5,811,238; 5,830,721; 5,834,252; and 5,837,458, and Patten et al., Curr. Opinion Biotechnol. 8:724-33 (1997); Harayama, Trends Biotechnol. 16(2):76-82 (1998); Hansson, et al., J. Mol. Biol. 287:265-76 (1999); and Lorenzo and Blasco, Biotechniques 24(2):308-13 (1998) (each of these patents and publications are hereby incorporated by reference in its entirety). In one embodiment, polynucleotides encoding antibodies of the invention may be altered by being subjected to random mutagenesis by error-prone PCR, random nucleotide insertion or other methods prior to recombination. In another embodiment, one or more portions of a polynucleotide encoding an antibody which portions

immunospecifically bind to BLyS may be recombined with one or more components, motifs, sections, parts, domains, fragments, etc. of one or more heterologous molecules.

[0341] Moreover, the antibodies of the present invention (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), can be fused to marker sequences, such as a polypeptides to facilitate purification. In preferred embodiments, the marker amino acid sequence is a hexahistidine polypeptide, such as the tag provided in a pQE vector (QIAGEN, Inc., 9259 Eton Avenue, Chatsworth, CA, 91311), among others, many of which are commercially available. As described in Gentz et al., Proc. Natl. Acad. Sci. USA 86:821-824 (1989), for instance, hexa-histidine provides for convenient purification of the fusion protein. Other peptide tags useful for purification include, but are not limited to, the hemagglutinin "HA" tag, which corresponds to an epitope derived from the influenza hemagglutinin protein (Wilson et al., Cell 37:767 (1984)) and the "flag" tag (DYKDDDDK, (SEQ ID No: 3238) Stratagene, La Jolla, CA).

The present invention further encompasses antibodies (including scFvs and [0342] other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), conjugated to a diagnostic or the apeutic agent. The antibodies can be used diagnostically to, for example, monitor or prognose the development or progression of a tumor as part of a clinical testing procedure to, e.g., determine the efficacy of a given treatment regimen. Detection can be facilitated by coupling the antibody to a detectable. substance. Examples of detectable substances include, but are not limited to, various enzymes, prosthetic groups, fluorescent materials, luminescent materials, bioluminescent materials, radioactive materials, positron emitting metals using various positron emission tomographies, and nonradioactive paramagnetic metal ions. The detectable substance may be coupled or conjugated either directly to the antibody or indirectly, through an intermediate (such as, for example, a linker known in the art) using techniques known in the art. See, for example, U.S. Patent No. 4,741,900 for metal ions which can be conjugated to antibodies for use as diagnostics according to the present invention. Examples of suitable enzymes include, but are not limited to, horseradish peroxidase, alkaline phosphatase, beta-galactosidase, or acetylcholinesterase; examples of suitable prosthetic group complexes include, but are not limited to, streptavidinlbiotin and avidin/biotin; examples of suitable fluorescent materials include, but are not limited to,

umbelliferone, fluorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine fluorescein, dansyl chloride or phycoerythrin; an example of a luminescent material includes, but is not limited to, luminol; examples of bioluminescent materials include, but are not limited to, luciferase, luciferin, and aequorin; and examples of suitable radioactive material include, but are not limited to, iodine (131 I, 125 I, 123 I, 121 I), carbon (14C), sulfur (35S). tritium (3H), indium (115mIn, 113mIn, 111In), and technetium (9Tc, 99mTc), thallium (201Ti), gallium (68Ga, 67Ga), palladium (109Pd), molybdenum (99Mo), xenon (133Xe). fluorine (18F), 153Sm, 177Lu, 159Gd, 149Pm, 140La, 175Yb, 166Ho, 90Y, 47Sc, 186Re, 188Re, 142Pr. ¹⁰⁵Rh, ⁹⁷Ru, ⁶⁸Ge, ⁵⁷Co, ⁶⁵Zn, ⁸⁵Sr, ³²P, ¹⁵³Gd, ¹⁶⁹Yb, ⁵¹Cr, ⁵⁴Mn, ⁷⁵Se, ¹¹³Sn, and ¹¹⁷Tin. [0343] Further, an antibody of the invention (including an scFy or other molecule comprising, or alternatively consisting of, antibody fragments or variants thereof), may be conjugated to a therapeutic moiety such as a cytotoxin, e.g., a cytostatic or cytocidal agent, a therapeutic agent or a radioactive metal ion, e.g., alpha-emitters such as, for example, ²¹³Bi. In specific embodiments, antibodies of the invention are attached to macrocyclic chelators useful for conjugating radiometal ions, including but not limited to. 111 In. 177 Lu. ⁹⁰Y, ¹⁶⁶Ho, and ¹⁵³Sm, to polypeptides. In preferred embodiments, the radiometal ion associated with the macrocyclic chelators attached to antibodies of the invention is 111 In. In preferred embodiments, the radiometal ion associated with the macrocyclic chelators attached to antibodies of the invention is 90Y. In specific embodiments, the macrocyclic chelator is 1,4,7,10-tetraazacyclododecane-N,N',N",N"'-tetraacetic acid (DOTA). In other specific embodiments, the DOTA is attached to the antibody of the invention via a linker molecule. Examples of linker molecules useful for conjugating DOTA to a polypeptide are commonly known in the art - see, for example, DeNardo et al., Clin Cancer Res. 4(10):2483-90, 1998; Peterson et al., Bioconjug. Chem. 10(4):553-7, 1999; and Zimmerman et al, Nucl. Med. Biol. 26(8):943-50, 1999 which are hereby incorporated by reference in their entirety.

[0344] A cytotoxin or cytotoxic agent includes any agent that is detrimental to cells and includes such molecules as small molecule toxins and enzymatically active

toxins of bacterial, fungal, plant, or animal origin, or fragments thereof. Examples include, but are not limited to, paclitaxol, cytochalasin B, gramicidin D, ethidium bromide, emetine, mitomycin, etoposide (VP-16), tenoposide, vincristine, vinblastine, colchicin, doxorubicin, daunorubicin, dihydroxy anthracin dione, mitoxantrone, mithramycin, actinomycin D, 1-dehydrotestosterone, glucocorticoids, procaine, tetracaine, lidocaine, propranolol, thymidine kinase, endonuclease, RNAse, and puromycin and frragments. variants or homologs thereof. Therapeutic agents include, but are not limited to. antimetabolites (e.g., methotrexate, 6-mercaptopurine, 6-thioguanine, cytarabine, 5fluorouracil decarbazine), alkylating agents (e.g., mechlorethamine, thioepa chlorambucil. melphalan, carmustine (BSNU) and lomustine (CCNU), cyclophosphamide, busulfan, dibromomannitol, streptozotocin, mitomycin C, and cisdichlorodiamine platinum (II) (DDP) cisplatin), anthracyclines (e.g., daunorubicin (formerly daunomycin) and doxorubicin), antibiotics (e.g., dactinomycin (formerly actinomycin), bleomycin, mithramycin, and anthramycin (AMC)), and anti-mitotic agents (e.g., vincristine and vinblastine), improsulfan, piposulfan, benzodopa, carboquone, meturedopa, uredopa, altretamine, triethylenemelamine, trietylenephosphoramide, triethylenethiophosphaoramide trimethylolomelamine, chlornaphazine, cholophosphamide, estramustine, ifosfamide, novembichin, phenesterine, prednimustine, trofosfamide, uracil mustard, chlorozotocin, fotemustine, nimustine, ranimustine, aclacinomysins, azaserine, cactinomycin, calichearnicin, carabicin, carminomycin, carzinophilin, chromomycins, detorubicin, 6-diazo-5-oxo-L-norleucine, epirubicin, esorubicin, idarubicin, marcellomycin, mycophenolic acid, nogalamycin, olivomycins, peplomycin, potfiromycin, quelamycin, rodorubicin, streptonigrin, tubercidin, ubenimex, zinostatin, zorubicin, denopterin, pteropterin, trimetrexate, fludarabine, thiamiprine, ancitabine, azacitidine, 6-azauridine, carmofur, dideoxyuridine, doxifluridine, enocitabine, floxuridine, 5-FU, calusterone, dromostanolone propionate, epitiostanol, mepitiostane, testolactone, aminoglutethimide, mitotane, trilostane, frolinic acid, aceglatone,

aldophosphamide glycoside, aminolevulinic acid, amsacrine, bestrabucil, bisantrene, edatraxate, defofamine, dernecolcine, diaziquone, elfornithine, elliptiniurn acetate, etoglucid, gallium nitrate, hydroxyurea, lentinan, lonidamine, mitoguazone, mopidamol, nitracrine, pentostatin, phenamet, pirarubicin, podophyllinic acid, 2-ethylhydrazide, procarbazine, PSKO, razoxane, sizofiran, spirogermanium, tenuazonic acid, triaziquone, 2, 2',2"-trichlorotriethylamine, urethan, vindesine, dacarbazine, mannomustine, mitobronitol, mitolactol, pipobroman, gacytosine, arabinoside ("Ara-C"), taxoids, e.g. paclitaxel (TAXOL", Bristol-Myers Squibb Oncology, Princeton, NJ) doxetaxel (TAXOTERE", Rhône-Poulenc Rorer, Antony, France), gemcitabine, ifosfamide, vinorelbine, navelbine, novantrone, teniposide, aminopterin, xeloda, ibandronate, CPT-I 1, topoisomerase inhibitor RFS 2000, difluoromethylornithine (DMFO), retinoic acid, esperamicins, capecitabine, and pharmaceutically acceptable salts, acids or derivatives of any of the above. Also included in this definition are anti-hormonal agents that act to regulate or inhibit hormone action on tumors such as anti-estrogens including for example tamoxifen, raloxifene, aromatase inhibiting 4(5)-imidazoles, 4 hydroxytamoxifen, trioxifene, keoxifene, LY 117018, onapristone, toremifene (Fareston), and anti-androgens such as flutamide, nilutamide, bicalutamide, leuprolide, and goserelin, and pharmaceutically acceptable salts, acids or derivatives of any of the above.

[0345] Techniques known in the art may be applied to label antibodies of the invention. Such techniques include, but are not limited to, the use of bifunctional conjugating agents (see e.g., U.S. Patent Nos. 5,756,065; 5,714,631; 5,696,239; 5,652,361; 5,505,931; 5,489,425; 5,435,990; 5,428,139; 5,342,604; 5,274,119; 4,994,560; and 5,808,003; the contents of each of which are hereby incorporated by reference in its entirety) and direct coupling reactions (e.g., Bolton-Hunter and Chloramine-T reaction).

[0346] The antibodies of the invention which are conjugates can be used for modifying a given biological response, the therapeutic agent or drug moiety is not to be construed as limited to classical chemical therapeutic agents. For example, the drug moiety may be a protein or polypeptide possessing a desired biological activity. Such

proteins may include, but are not limited to, for example, a toxin such as abrin, ricin A, alpha toxin, pseudomonas exotoxin, or diphtheria toxin, saporin, momordin, gelonin, pokeweed antiviral protein, alpha-sarcin and cholera toxin; a protein such as tumor necrosis factor, alpha-interferon, beta-interferon, nerve growth factor, platelet derived growth factor, tissue plasminogen activator, an apoptotic agent, e.g., TNF-alpha, TNF-beta, AIM I (see, International Publication No. WO 97/33899), AIM II (see, International Publication No. WO 97/34911), Fas Ligand (Takahashi et al., Int. Immunol., 6:1567-1574 (1994)), VEGI (see, International Publication No. WO 99/23105), a thrombotic agent or an anti-angiogenic agent, e.g., angiostatin or endostatin; or, biological response modifiers such as, for example, lymphokines, interleukin-1 (IL-1), interleukin-2 (IL-2), interleukin-6 (IL-6), granulocyte macrophage colony stimulating factor (GM-CSF), granulocyte colony stimulating factor (G-CSF), or other growth factors.

[0347] Antibodies of the invention (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), may also be attached to solid supports, which are particularly useful for immunoassays or purification of the target antigen. Such solid supports include, but are not limited to, glass, cellulose, polyacrylamide, nylon, polystyrene, polyvinyl chloride or polypropylene.

[0348] Techniques for conjugating a therapeutic moiety to antibodies are well known, see, e.g., Arnon et al., "Monoclonal Antibodies For Immunotargeting Of Drugs In Cancer Therapy", in Monoclonal Antibodies And Cancer Therapy, Reisfeld et al. (eds.), pp. 243-56 (Alan R. Liss, Inc. 1985); Hellstrom et al., "Antibodies For Drug Delivery", in Controlled Drug Delivery (2nd Ed.), Robinson et al. (eds.), pp. 623-53 (Marcel Dekker, Inc. 1987); Thorpe, "Antibody Carriers Of Cytotoxic Agents In Cancer Therapy: A Review", in Monoclonal Antibodies '84: Biological And Clinical Applications, Pinchera et al. (eds.), pp. 475-506 (1985); "Analysis, Results, And Future Prospective Of The Therapeutic Use Of Radiolabeled Antibody In Cancer Therapy", in Monoclonal Antibodies For Cancer Detection And Therapy, Baldwin et al. (eds.), pp. 303-16 (Academic Press 1985), and Thorpe et al., "The Preparation And Cytotoxic Properties Of Antibody-Toxin Conjugates", Immunol. Rev. 62:119-58 (1982).

[0349] Alternatively, an antibody of the invention can be conjugated to a second antibody to form an antibody heteroconjugate as described by Segal in U.S. Patent No. 4,676,980, which is incorporated herein by reference in its entirety.

[0350] An antibody of the invention (including an scFv or and other molecule comprising, or alternatively consisting of, an antibody fragment or variant thereof), with or without a therapeutic moiety conjugated to it, administered alone or in combination with cytotoxic factor(s) and/or cytokine(s) can be used as a therapeutic.

Use of Antibodies for Epitope Mapping

The present invention provides antibodies (including scFvs and other molecules comprising, or alternatively consisting of, antibody fragments or variants thereof), that can be used to identify epitopes of BLyS. In particular, the antibodies of the present invention can be used to identify epitopes of human BLyS (SEQ ID NOS:3228 and/or 3229) or BLyS expressed on human monocytes; murine BLyS (SEQ ID NOS:3230 and/or 3231) or BLyS expressed on murine monocytes; rat BLyS (either the soluble forms as given in SEQ ID NOS:3232, 3233, 3234 and/or 3235 or in a membrane associated form, e.g., on the surface of rat monocytes); or monkey BLyS (e.g., the monkey BLyS polypeptides of SEQ ID NOS:3236 and/or 3237, the soluble form of monkey BLyS, or BLyS expressed on monkey monocytes)using techniques described herein or otherwise known in the art. Fragments which function as epitopes may be produced by any conventional means. (See, e.g., Houghten, Proc. Natl. Acad. Sci. USA 82:5131-5135 (1985), further described in U.S. Patent No. 4,631,211.)

Diagnostic Uses of Antibodies

[0352] Labeled antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor diseases and/or disorders associated with the aberrant expression and/or activity of BLyS or BLyS receptor. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample from an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase or decrease in the assayed level of BLyS compared to the standard level of BLyS is indicative of aberrant expression.

[0353] By "biological sample" is intended any fluids and/or cells obtained from an individual, body fluid, body tissue, body cell, cell line, tissue culture, or other source which may contain BLyS protein or mRNA. Body fluids include, but are not limited to, sera, plasma, urine, synovial fluid, spinal fluid, saliva, and mucous. Tissues samples may be taken from virtually any tissue in the body. Tissue samples may also be obtained from autopsy material. Methods for obtaining tissue biopsies and body fluids from mammals are well known in the art. Where the biological sample is to include mRNA, a tissue biopsy is the preferred source.

[0354] The invention also provides for the detection of aberrant expression of BLyS receptor comprising (a) assaying the expression of BLyS receptor in a biological sample from an individual using one or more antibodies or fragments or variants thereof that immunospecifically binds only to soluble BLyS, but does not inhibit BLyS /BLyS receptor binding. Such an antibody, by way of an example that is not to be construed as limiting, would be one that is able to capture a biotinylated BLyS from solution (see Example 8), but that would not prevent BLyS from binding to IM-9 cells (see Example 3). and (b) comparing the level of BLyS receptor with a standard level of BLyS receptor, e.g., in normal tissue or cell samples, whereby an increase or decrease in the assayed level of BLyS receptor compared to the standard level of BLyS receptor is indicative of aberrant expression.

[0355] Antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor autoimmune disorders and/or immunodeficiencies, and/or diseases or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample from an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase or decrease in the assayed level of BLyS compared to the standard level of BLyS is indicative of an autoimmune disorder or disease and/or an immunodeficiency. In specific embodiments, an increase in the assayed level of BLyS is indicative of an autoimmune disorder or disease. In other specific embodiments, a decrease in the assayed level of BLyS is indicative of an immunodeficiency.

[0356] Antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) which specifically bind to BLyS but, do not inhibit BLyS/BLyS receptor binding can be used for diagnostic purposes to detect, diagnose, prognose, or monitor autoimmune disorders and/or immunodeficiencies, and/or diseases or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS receptor comprising: (a) assaying the expression of BLyS receptor in a biological sample from an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS receptor with a standard level of BLyS receptor, e.g., in normal biological samples, whereby an increase or decrease in the assayed level of BLyS receptor compared to the standard level of BLyS receptor is indicative of an autoimmune disorder or disease and/or an immunodeficiency. In specific embodiments, an increase in the assayed level of BLyS receptor is indicative of an autoimmune disorder or disease. In other specific embodiments, a decrease in the assayed level of BLyS receptor is indicative of an immunodeficiency.

Autoimmune disorders, diseases, or conditions that may be detected, [0357] diagnosed, prognosed, or monitored using the antibodies of the invention include, but are not limited to, autoimmune hemolytic anemia, autoimmune neonatal thrombocytopenia, idiopathic thrombocytopenia purpura, autoimmune neutropenia, autoimmunocytopenia, hemolytic anemia, antiphospholipid syndrome, dermatitis, gluten-sensitive enteropathy, allergic encephalomyelitis, myocarditis, relapsing polychondritis, rheumatic heart disease, glomerulonephritis (e.g., IgA nephropathy), Multiple Sclerosis, Neuritis, Uveitis Ophthalmia, Polyendocrinopathies, Purpura (e.g., Henloch-Scoenlein purpura), Reiter's Disease, Stiff-Man Syndrome, Autoimmune Pulmonary Inflammation, myocarditis, IgA glomerulonephritis, dense deposit disease, rheumatic heart disease, Guillain-Barre Syndrome, diabetes mellitus (e.g. Type I diabetes mellitus or insulin dependent diabetes mellitis), juvenile onset diabetes, and autoimmune inflammatory eye, autoimmune thyroiditis, hypothyroidism (i.e., Hashimoto's thyroiditis, systemic lupus erhythematosus, discoid lupus, Goodpasture's syndrome, Pemphigus, Receptor autoimmunities such as, for example, (a) Graves' Disease, (b) Myasthenia Gravis, and (c) insulin resistance, autoimmune hemolytic anemia, autoimmune thrombocytopenic purpura, rheumatoid arthritis, schleroderma with anti-collagen antibodies, mixed connective tissue disease,

polymyositis/dermatomyositis, pernicious anemia (Addison's disease), idiopathic Addison's disease, infertility, glomerulonephritis such as primary glomerulonephritis and IgA nephropathy, bullous pemphigoid, Sjögren's syndrome, diabetes millitus, and adrenergic drug resistance (including adrenergic drug resistance with asthma or cystic fibrosis), chronic active hepatitis, primary biliary cirrhosis, other endocrine gland failure, vitiligo, vasculitis, post-MI, cardiotomy syndrome, urticaria, atopic dermatitis, asthma, inflammatory myopathies, and other inflammatory, granulomatous, degenerative, and atrophic disorders and other disorders such as inflammatory skin diseases including psoriasis and sclerosis, responses associated with inflammatory bowel disease (such as Crohn's disease and ulcerative colitis), respiratory distress syndrome (including adult respiratory distress syndrome, ARDS), meningitis, encephalitis, colitis, allergic conditions such as eczema and other conditions involving infiltration of T cells and chronic inflammatory responses, atherosclerosis, leukocyte adhesion deficiency, Reynaud's syndrome, and immune responses associated with acute and delayed hypersensitivity mediated by cytokines and T-lymphocytes typically found in tuberculosis, sarcoidosis, granulomatosis and diseases involving leukocyte diapedesis, central nervous system (CNS) inflammatory disorder, multiple organ injury syndrome, antigen-antibody complex mediated diseases, anti-glomerular basement membrane disease, Lambert-Eaton myasthenic syndrome, Beheet disease, giant cell arteritis, immune complex nephritis, IgA nephropathy, IgM polyneuropathies or autoimmune thrombocytopenia etc.

[0358] In specific embodiments, the present invention encompasses methods and compositions for detecting, diagnosing and/or prognosing diseases or disorders associated with hypergammaglobulinemia (e.g., AIDS, autoimmune diseases, and some immunodeficiencies). In other specific embodiments, the present invention encompasses methods and compositions for detecting, diagnosing and/or prognosing diseases or disorders associated with hypogammaglobulinemia (e.g., an immunodeficiency).

[0359] Immunodeficiencies that may be detected, diagnosed, prognosed, or monitored using the antibodies of the invention include, but are not limited to, severe combined immunodeficiency (SCID)-X linked, SCID-autosomal, adenosine deaminase

deficiency (ADA deficiency), X-linked agammaglobulinemia (XLA), Bruton's disease, congenital agammaglobulinemia, X-linked infantile agammaglobulinemia, acquired agammaglobulinemia, adult onset agammaglobulinemia, late-onset agammaglobulinemia, dysgammaglobulinemia, hypogammaglobulinemia, transient hypogammaglobulinemia of infancy, unspecified hypogammaglobulinemia, agammaglobulinemia, common variable immunodeficiency (CVID) (acquired), Wiskott-Aldrich Syndrome (WAS), X-linked immunodeficiency with hyper IgM, non X-linked immunodeficiency with hyper IgM, selective IgA deficiency, IgG subclass deficiency (with or without IgA deficiency), antibody deficiency with normal or elevated Igs, immunodeficiency with thymoma, Ig heavy chain deletions, kappa chain deficiency, B cell lymphoproliferative disorder (BLPD), selective IgM immunodeficiency, recessive agammaglobulinemia (Swiss type), reticular dysgenesis, neonatal neutropenia, severe congenital leukopenia, thymic alymphoplasia-aplasia or dysplasia with immunodeficiency, ataxia-telangiectasia, short limbed dwarfism, X-linked lymphoproliferative syndrome (XLP), Nezelof syndromecombined immunodeficiency with Igs, purine nucleoside phosphorylase deficiency (PNP), MHC Class II deficiency (Bare Lymphocyte Syndrome) and severe combined immunodeficiency.

[0360] Elevated levels of soluble BLyS have been observed in the serum of patients with Systemic Lupus Erythematosus (SLE). In comparing the sera of 150 SLE patients with that of 38 control individuals, it was found that most of the SLE patients had more than 5ng/ml of serum BLyS, more than 30% of SLE patients had levels greater than 10ng/ml, and approximately 10% of SLE patients had serum BLyS levels greater than 20ng/ml. In contrast, the majority of normal controls had BLyS levels less than 5ng/ml, and less than 10% had levels higher than 10ng/ml. The elevated levels of BLyS protein in sera is present in the soluble form and has biologic activity as assayed by the ability to stimulate anti-IgM treated B cells in vitro. SLE patients with more than 15ng/ml serum BLyS were also found to have elevated levels of anti-dsDNA antibodies compared to both normal controls and SLE patients with less than 5ng/ml of serum BLyS.(unpublished data).

[0361] In addition the serum of two subgroups of patients which were positive for anti-nuclear antibodies (ANA+) but did not meet the formal requirements of the American College of Rheumatology (ACR) for classification of SLE were analyzed for BLyS levels.

The first subgroup of sera was ANA+ sera that came from patients who did not present with the clinical impression of SLE. This group had only slightly elevated levels of BLyS (~9ng/ml BLyS). The second subgroup however, which was ANA+ sera from patients who presented with the clinical impression of SLE, had significantly increased BLyS levels (~15ng/ml). These results suggest that an elevated level of BLyS precedes the formal fulfillment of the ACR criteria. The ACR criteria are described in Tan, E.M., et al, Arthritis and Rheumatism 25:1271 – 1277 (1982).

[0362] Thus in specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor Systemic Lupus Erythematosus or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of SLE.

[0363] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor IgA nephropathy or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of IgA nephropathy.

[0364] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor Sjögren's Syndrome or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby

an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of Sjögren's Syndrome.

[0365] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor HIV infection or conditions associated therewith (e.g. AIDS). The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of HIV infection.

[0366] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor Myasthenia Gravis or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of Myasthenia Gravis.

[0367] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor idiopathic thrombocytopenic purpura (ITP) or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of idiopathic thrombocytopenic purpura (ITP).

[0368] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor hemolytic anemia or conditions associated therewith. The invention

provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of hemolytic anemia.

[0369] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor thyroiditis or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of thyroiditis.

[0370] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor Goodpasture's syndrome or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of Goodpasture's syndrome.

[0371] In other specific embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor multiple sclerosis or conditions associated therewith. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an

increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of multiple sclerosis.

[0372] In additional embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor Rheumatoid Arthritis. The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample (e.g., serum and synovial fluid) of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of Rheumatoid arthritis.

[0373] In additional embodiments, antibodies of the invention which specifically bind to BLyS can be used for diagnostic purposes to detect, diagnose, prognose, or monitor an immune-based rheumatologic disease, (e.g., SLE, rheumatoid arthritis, CREST syndrome (a variant of scleroderma characterized by calcinosis, Raynaud's phenomenon, esophageal motility disorders, sclerodactyly, and telangiectasia.), Seronegative spondyloarthropathy (SpA), Polymyositis/dermatomyositis, Microscopic polyangiitis, Hepatitis C-associated arthritis, Takayasu's arteritis, and undifferentiated connective tissue disorder). The invention provides for the detection of aberrant expression of BLyS comprising: (a) assaying the expression of BLyS in a biological sample (e.g., serum and synovial fluid) of an individual using one or more antibodies of the invention that immunospecifically binds to BLyS; and (b) comparing the level of BLyS with a standard level of BLyS, e.g., in normal biological samples, whereby an increase in the assayed level of BLyS compared to the standard level of BLyS is indicative of monitor an immune-based rheumatologic disease.

It has been observed, that serum BLyS levels inversely correlate with nephrotic range proteinuria (>3gm proteinuria in a 24 hour urine collection) using a sample of 71 SLE patients (p=0.019). Proteinuria was determined in 71 SLE patients within one month of phlebotomy for serum BLyS determination. Serum BLyS was classified as low, normal, or high based on the 5th through 95th percentiles for normal controls. Nephrotic-range proteinuria was inversely correlated with serum Neutrokinealpha levels. Thus, in specific embodiments, serum levels of BLyS (determined using one

or more antibodies of the present invention) in individuals diagnosed with an immune based rheumatologic disease (e.g., SLE, rheumatoid arthritis, CREST syndrome (a variant of scleroderma characterized by calcinosis, Raynaud's phenomenon, esophageal motility disorders, sclerodactyly, and telangiectasia.), seronegative spondyloarthropathy (SpA), polymyositis/dermatomyositis, microscopic polyangiitis, hepatitis C-asociated arthritis, Takayasu's arteritis, and undifferentiated connective tissue disorder) may be used to determine, diagnose, prognose, or monitor the severity of certain aspects or symptoms of the disease, such as nephrotic-range proteinuria.

[0375] In another specific embodiment, antibodies of the invention are used to diagnose, prognose, treat, or prevent conditions associated with CVID, including, but not limited to, conditions associated with acute and recurring infections (e.g., pneumonia, bronchitis, sinusitis, otitis media, sepsis, meningitis, septic arthritis, and osteomyelitis), chronic lung disease, autoimmunity, granulomatous disease, lymphoma, cancers (e.g., cancers of the breast, stomach, colon, mouth, prostate, lung, vagina, ovary, skin, and melanin forming cells (i.e. melanoma), inflammatory bowel disease (e.g., Crohn's disease, ulcerative colitis, and ulcerative proctitis), malabsorption, Hodgkin's disease, and Waldenstrom's macroglobulinemia.

[0376] The invention provides a diagnostic assay for diagnosing or prognosing a disease or disorder, comprising: (a) assaying for the level of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically bind to BLyS; and (b) comparing the level of BLyS with a standard BLyS level, e.g., in a biological sample from a patient without the disease or disorder, whereby an increase or decrease in the assayed BLyS level compared to the standard level of BLyS is indicative of a particular disease or disorder. With respect to cancer, the presence of a relatively high amount of BLyS in biopsied tissue from an individual may indicate a predisposition for the development of the disease, or may provide a means for detecting the disease prior to the appearance of actual clinical symptoms. A more definitive diagnosis of this type may allow health professionals to employ preventative measures or aggressive treatment earlier thereby preventing the development or further progression of the cancer.

[0377] In specific embodiments, the presence of a relatively high amount of

membrane-bound BLyS in a biological sample is indicative of monocytic cell related leukemias or lymphomas, such as, for example acute myelogenous leukemia and/or the severity thereof.

[0378] In other specific embodiments, the presence of a relatively high amount of BLyS receptor in a biological sample (as determined using antibodies of the invention that bind to soluble BLyS, but do not inhibit BLyS/BLyS receptor binding) is indicative of B cell related leukemias or lymphomas (e.g., chronic lymphocytic leukemia, multiple myeloma, non-Hodgkin's lymphoma, and Hodgkin's disease), and/or the severity thereof.

[0379] In specific embodiments, the invention provides a diagnostic assay for diagnosing or prognosing Systemic Lupus Erythematosus, comprising: (a) assaying for the level of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically bind to BLyS; and (b) comparing the level of BLyS with a standard BLyS level, e.g., in a biological sample from a patient without Systemic Lupus Erythematosus, whereby an increase in the assayed BLyS level compared to the standard level of BLyS is indicative of Systemic Lupus Erythematosus.

[0380] In specific embodiments, the invention provides a diagnostic assay for diagnosing or prognosing a Rheumatoid Arthritis, comprising: (a) assaying for the level of BLyS in a biological sample of an individual using one or more antibodies of the invention that immunospecifically bind to BLyS; and (b) comparing the level of BLyS with a standard BLyS level, e.g., in a biological sample from a patient without Rheumatoid Arthritis, whereby an increase or decrease in the assayed BLyS level compared to the standard level of BLyS is indicative of Rheumatoid Arthritis.

[0381] The invention provides a diagnostic assay for diagnosing or prognosing a disease or disorder, comprising: (a) assaying for the level of BLyS receptor in cells or a tissue sample of an individual using one or more antibodies of the invention that immunospecifically binds only to soluble BLyS, but does not neutralize BLyS/BLyS receptor binding; and (b) comparing the level of BLyS receptor with a standard BLyS receptor level, e.g., in a tissue sample from a patient without the disease or disorder, whereby an increase or decrease in the assayed BLyS receptor level compared to the standard level of BLyS receptor is indicative of a particular disease or disorder. With respect to cancer, the presence of a relatively high amount of BLyS receptor in biopsied tissue from an individual may indicate a predisposition for the development of the disease,

or may provide a means for detecting the disease prior to the appearance of actual clinical symptoms. A more definitive diagnosis of this type may allow health professionals to employ preventative measures or aggressive treatment earlier thereby preventing the development or further progression of the cancer.

Antibodies of the invention (including molecules comprising, or [0382] alternatively consisting of, antibody fragments or variants thereof) can be used to assay protein levels in a biological sample using classical immunohistological methods as described herein or as known to those of skill in the art (e.g., see Jalkanen, et al., J. Cell. Biol. 101:976-985 (1985); Jalkanen, et al., J. Cell. Biol. 105:3087-3096 (1987)). Other antibody-based methods useful for detecting protein gene expression include immunoassays, such as the enzyme linked immunosorbent assay (ELISA) and the radioimmunoassay (RIA). Suitable antibody assay labels are known in the art and include enzyme labels, such as, glucose oxidase, alkaline phosphatase, and horseradish peroxidase; radioisotopes, such as iodine (121I, 123I, 125I, 131I), carbon (14C), sulfur (35S), tritium (³H), indium (¹¹¹In, ¹¹²In, ^{113m}In, ^{115m}In), technetium (⁹⁹Tc, ^{99m}Tc), thallium (²⁰¹Ti), gallium (68Ga, 67Ga), palladium (108Pd), molybdenum (99Mo), xenon (133Xe), fluorine (18F), ¹⁵³Sm, ¹⁷⁷Lu, ¹⁵⁹Gd, ¹⁴⁹Pm, ¹⁴⁰La, ¹⁷⁵Yb, ¹⁶⁶Ho, ⁹⁰Y, ⁴⁷Sc, ¹⁸⁶Re, ¹⁸⁸Re, ¹⁴²Pr, ¹⁰⁵Rh, and ⁹⁷Ru; luminescent labels, such as luminol; and fluorescent labels, such as fluorescein and rhodamine, and biotin.

[0383] One aspect of the invention is the detection and diagnosis of a disease or disorder associated with aberrant expression of BLyS or BLyS receptor in an animal, preferably a mammal and most preferably a human. In one embodiment, diagnosis comprises: a) administering (for example, parenterally, subcutaneously, or intraperitoneally) to a subject an effective amount of a labeled antibody of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically binds to BLyS; b) waiting for a time interval following the administering for permitting the labeled antibody to preferentially concentrate at sites in the subject where BLyS is expressed (and for unbound labeled molecule to be cleared to background level); c) determining background level; and d) detecting the labeled antibody in the subject, such that detection of labeled antibody or fragment thereof above the background level and above or below the level observed in a person without the disease or disorder indicates that the subject has a particular disease or

disorder associated with aberrant expression of BLyS or BLyS receptor. Background level can be determined by various methods including, comparing the amount of labeled molecule detected to a standard value previously determined for a particular system.

It will be understood in the art that the size of the subject and the imaging system used will determine the quantity of imaging moiety needed to produce diagnostic images. In the case of a radioisotope moiety, for a human subject, the quantity of radioactivity injected will normally range from about 5 to 20 millicuries of ⁹⁹Tc. The labeled antibody will then preferentially accumulate at the location of cells which contain the specific protein. *In vivo* tumor imaging is described in S.W. Burchiel *et al.*, "Immunopharmacokinetics of Radiolabeled Antibodies and Their Fragments." (Chapter 13 in Tumor Imaging: The Radiochemical Detection of Cancer, S.W. Burchiel and B. A. Rhodes, eds., Masson Publishing Inc. (1982).

[0385] Depending on several variables, including the type of label used and the mode of administration, the time interval following the administration for permitting the labeled molecule to preferentially concentrate at sites in the subject and for unbound labeled molecule to be cleared to background level is 6 to 48 hours or 6 to 24 hours or 6 to 12 hours. In another embodiment the time interval following administration is 5 to 20 days or 5 to 10 days.

[0386] In an embodiment, monitoring of the disease or disorder is carried out by repeating the method for diagnosing the disease or disorder, for example, one month after initial diagnosis, six months after initial diagnosis, one year after initial diagnosis, etc.

[0387] Presence of the labeled molecule can be detected in the patient using methods known in the art for *in vivo* scanning. These methods depend upon the type of label used. Skilled artisans will be able to determine the appropriate method for detecting a particular label. Methods and devices that may be used in the diagnostic methods of the invention include, but are not limited to, computed tomography (CT), whole body scan such as position emission tomography (PET), magnetic resonance imaging (MRI), and sonography.

[0388] In a specific embodiment, the molecule is labeled with a radioisotope and is detected in the patient using a radiation responsive surgical instrument (Thurston et al., U.S. Patent No. 5,441,050). In another embodiment, the molecule is labeled with a fluorescent compound and is detected in the patient using a fluorescence responsive

scanning instrument. In another embodiment, the molecule is labeled with a positron emitting metal and is detected in the patient using positron emission-tomography. In yet another embodiment, the molecule is labeled with a paramagnetic label and is detected in a patient using magnetic resonance imaging (MRI).

Immunophenotyping

[0389] The antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may be utilized for immunophenotyping of cell lines and biological samples by their BLyS expression or BLyS receptor expression. Various techniques can be utilized using antibodies, fragments, or variants of the invention to screen for cellular populations (i.e., immune cells, particularly monocytic cells or B-cells) expressing BLyS or BLyS receptor, and include magnetic separation using antibody-coated magnetic beads, "panning" with antibody attached to a solid matrix (i.e., plate), and flow cytometry (see, e.g., U.S. Patent 5,985,660; and Morrison et al., Cell, 96:737-49 (1999)).

[0390] These techniques allow for the screening of particular populations of cells, such as might be found with hematological malignancies (i.e., minimal residual disease (MRD) in acute leukemic patients) and "non-self" cells in transplantations to prevent Graft-versus-Host Disease (GVHD). Alternatively, these techniques allow for the screening of hematopoietic stem and progenitor cells capable of undergoing proliferation and/or differentiation, as might be found in human umbilical cord blood.

[0391] In one embodiment, antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) are used to identify cells of monocytic or B cell origin.

Therapeutic Uses of Antibodies

[0392] The present invention is further directed to antibody-based therapies which involve administering antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) to an animal, preferably a mammal, and most preferably a human, patient for treating one or more of the disclosed diseases, disorders, or conditions. Therapeutic compounds of the invention include, but are not limited to, antibodies of the invention and nucleic acids encoding antibodies (and

anti-idiotypic antibodies) of the invention as described herein. The antibodies of the invention can be used to treat, ameliorate or prevent diseases, disorders or conditions associated with aberrant expression and/or activity of BLyS or BLyS receptor, including, but not limited to, any one or more of the diseases, disorders, or conditions described herein. The treatment and/or prevention of diseases, disorders, or conditions associated with aberrant BLyS expression and/or activity or aberrant BLyS receptor expression and/or activity includes, but is not limited to, alleviating symptoms associated with those diseases, disorders or conditions. Antibodies of the invention may be provided in pharmaceutically acceptable compositions as known in the art or as described herein.

Antibodies of the present invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that function as agonists or antagonists of BLyS, preferably of BLyS-induced signal transduction, can be administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, lack of BLyS function, aberrant BLyS receptor expression, or lack of BLyS receptor function. For example, antibodies of the invention which disrupt the interaction between BLyS and its receptor may be administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, excessive BLyS function, aberrant BLyS receptor expression, or excessive of BLyS receptor function. Antibodies of the invention which do not prevent BLyS from binding its receptor but inhibit or downregulate BLyS-induced signal transduction can be administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, excessive BLyS function, aberrant BLyS receptor expression, or excessive BLyS receptor function. In particular, antibodies of the present invention which prevent BLyS-induced signal transduction by specifically recognizing the unbound BLyS, receptor-bound BLyS or both unbound and receptor-bound BLyS can be administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, excessive BLyS function, aberrant BLyS receptor expression, or excessive BLyS receptor function. The ability of an antibody of the invention to inhibit or downregulate BLyS-induced signal transduction may be determined by techniques described herein or otherwise known in the art. For example, BLySinduced receptor activation and the activation of signaling molecules can be determined by detecting the phosphorylation (e.g., tyrosine or serine/threonine) of the receptor or a

signaling molecule by immunoprecipitation followed by western blot analysis (for example, as described herein).

In a specific embodiment, an antibody of the present invention (including [0394] molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that inhibits or downregulates BLyS activity by at least 95%, at least 90%, at least 85%, at least 80%, at least 75%, at least 70%, at least 60%, at least 50%, at least 45%, at least 40%, at least 45%, at least 35%, at least 30%, at least 25%, at least 20%, or at least 10% relative to BLyS activity in absence of the antibody is administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, excessive BLyS function, aberrant BLyS receptor expression, or excessive BLyS receptor function. In another embodiment, a combination of antibodies, a combination of antibody fragments, a combination of antibody variants, or a combination of antibodies, antibody fragments, and/or variants that inhibit or downregulate BLyS activity by at least 95%, at least 90%, at least 85%, at least 80%, at least 75%, at least 70%, at least 65%, at least 60%, at least 55%, at least 50%, at least 45%, at least 40%, at least 45%, at least 35%, at least 30%, at least 25%, at least 20%, or at least 10% relative to BLyS activity in absence of said antibodies, antibody fragments, and/or antibody variants are administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, excessive BLyS function, aberrant BLyS receptor expression, or excessive BLyS receptor function.

[0395] Further, antibodies of the present invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) which activate BLyS-induced signal transduction can be administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, lack of BLyS function, aberrant BLyS receptor expression, or lack of BLyS receptor function. These antibodies may potentiate or activate either all or a subset of the biological activities of BLyS-mediated receptor activation, for example, by inducing multimerization of BLyS and/or multimerization of the receptor. The antibodies of the invention may be administered with or without being pre-complexed with BLyS. In a specific embodiment, an antibody of the present invention that increases BLyS activity by at least 5%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 40%, at least 50%, at least 50%, at least 70%, at least 70%, at least 50%, at least 50%,

75%, at least 80%, at least 85%, at least 90%, at least 95%, or at least 99% relative to BLyS activity in absence of the antibody is administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression, lack of BLyS function, aberrant BLyS receptor expression, or lack of BLyS receptor function. In another embodiment, a combination of antibodies, a combination of antibody fragments, a combination of antibody variants, or a combination of antibodies, antibody fragments and/or antibody variants that increase BLyS activity by at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 65%, at least 70%, at least 75%, at least 80%, at least 90%, at least 99% relative to BLyS activity in absence of the said antibodies or antibody fragments and/or antibody variants is administered to an animal to treat, prevent or ameliorate a disease or disorder associated with aberrant BLyS expression or lack of BLyS function or aberrant BLyS receptor expression or lack of BLyS receptor function.

[0396] One or more antibodies of the present invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS may be used locally or systemically in the body as a therapeutic. The antibodies of this invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may also be advantageously utilized in combination with other monoclonal or chimeric antibodies, or with lymphokines or hematopoietic growth factors (such as, e.g., IL-2, IL-3 and IL-7), for example, which serve to increase the number or activity of effector cells which interact with the antibodies.

[0397] The antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) may be administered alone or in combination with other types of treatments (e.g., radiation therapy, chemotherapy, hormonal therapy, immunotherapy, anti-tumor agents, anti-angiogenesis and anti-inflammatory agents). Generally, administration of products of a species origin or species reactivity (in the case of antibodies) that is the same species as that of the patient is preferred. Thus, in a preferred embodiment, human antibodies, fragments, or variants, (e.g., derivatives), or nucleic acids, are administered to a human patient for therapy or prophylaxis.

It is preferred to use high affinity and/or potent in vivo inhibiting and/or [0398] neutralizing antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) that immunospecifically bind to BLyS, or polynucleotides encoding antibodies that immunospecifically bind to BLyS, for both immunoassays directed to and therapy of disorders related to BLyS polynucleotides or polypeptides, including fragments thereof. Such antibodies will preferably have an affinity for BLyS and/or BLyS fragments. Preferred binding affinities include those with a dissociation constant or K_D less than or equal to 5 X 10⁻² M, 10⁻² M, 5 X 10⁻³ M, 10⁻³ M, 5 X 10⁻⁴ M, 10⁻⁴ M, 5 X 10⁻⁵ M, or 10⁻⁵ M. More preferably, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with a dissociation constant or K_D less than or equal to 5 X 10⁻⁶ M, 10⁻⁶ M, 5 X 10⁻⁷ M, 10⁻⁷ M, 5 X 10⁻⁸ M, or 10⁻⁸ M. Even more preferably, antibodies of the invention bind BLyS polypeptides or fragments or variants thereof with a dissociation constant or K_D less than or equal to 5 X 10⁻⁹ M, 10⁻⁹ M, $5 \times 10^{-10} \text{ M}, 10^{-10} \text{ M}, 5 \times 10^{-11} \text{ M}, 10^{-11} \text{ M}, 5 \times 10^{-12} \text{ M}, 10^{-12} \text{ M}, 5 \times ^{-13} \text{ M}, 10^{-13} \text{ M}, 5 \times ^{-13} \text{ M}$ 10⁻¹⁴ M, 10⁻¹⁴ M, 5 X 10⁻¹⁵ M, or 10⁻¹⁵ M. The invention encompasses antibodies that bind BLyS polypeptides with a dissociation constant or KD that is within any one of the ranges that are between each of the individual recited values.

[0399] In a preferred embodiment, antibodies of the invention neutralize BLyS activity. In another preferred embodiment, antibodies of the invention inhibit B cell proliferation.

[0400] In a preferred embodiment, antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) inhibit or reduce binding of the soluble form of BLyS to a BLyS receptor. In another preferred embodiment antibodies of the invention inhibit or reduce B cell proliferation induced by the soluble form of BLyS. In another preferred embodiment antibodies of the invention inhibit or reduce immunoglobulin production induced by the soluble form of BLyS.

[0401] In a preferred embodiment, antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) inhibit or reduce binding of membrane-bound BLyS to a BLyS receptor. In another preferred embodiment, antibodies of the invention inhibit or reduce B cell proliferation induced by the membrane-bound form of BLyS. In another preferred

embodiment, antibodies of the invention inhibit or reduce immunoglobulin production induced by the membrane bound form of BLyS.

[0402] In a preferred embodiment, antibodies of the invention (including molecules comprising, or alternatively consisting of, antibody fragments or variants thereof) inhibit or reduce binding of both the soluble and membrane-bound forms of BLyS to a BLyS receptor. In another preferred embodiment, antibodies of the invention inhibit or reduce B cell proliferation induced by either or both forms of BLyS. In another preferred embodiment, antibodies of the invention inhibit or reduce immunoglobulin production induced by either or both forms of BLyS.

[0403] In one embodiment, the invention provides a method of delivering antibody conjugates of the invention to targeted cells, such as, for example, monocytic cells expressing the membrane-bound form of BLyS, or B cells expressing a BLyS receptor.

[0404] In one embodiment, the invention provides a method for the specific delivery of antibodies and antibody conjugates of the invention to cells by administering molecules of the invention that are associated with heterologous polypeptides or nucleic acids. In one example, the invention provides a method for delivering a therapeutic protein into the targeted cell. In another example, the invention provides a method for delivering a single stranded nucleic acid (e.g., antisense or ribozymes) or double stranded nucleic acid (e.g., DNA that can integrate into the cell's genome or replicate episomally and that can be transcribed) into the targeted cell.

[0405] In another embodiment, the invention provides a method for the specific destruction of cells (e.g., the destruction of tumor cells) by administering antibodies or antibody conjugates of the invention (e.g., antibodies conjugated with radioisotopes, toxins, or cytotoxic prodrugs). In a specific embodiment, the invention provides a method for the specific destruction of cells of monocytic lineage (e.g., monocytic cell related leukemias or lymphomas, such as, for example acute myelogenous leukemia) by administering antibodies or antibody conjugates of the invention (e.g., antibodies conjugated with radioisotopes, toxins, or cytotoxic prodrugs) that immunospecifically bind the membrane-bound form of BLyS. In another specific embodiment, the invention provides a method for the specific destruction of cells of B cell lineage (e.g., B cell related leukemias or lymphomas (e.g., chronic lymphocytic leukemia, multiple myeloma, non-Hodgkin's lymphoma, and Hodgkin's disease) by administering antibodies or antibody

conjugates of the invention (e.g., antibodies conjugated with radioisotopes, toxins, or cytotoxic prodrugs) that bind soluble BLyS, but do not inhibit BLyS binding to a BLyS receptor on B cells.

[0406] In another preferred embodiment antibodies of the invention (including antibody fragments and variants) promote or enhance B cell proliferation induced by the soluble form of BLyS. In another preferred embodiment, antibodies of the invention (including antibody fragments and variants) promote or enhance B cell proliferation induced by the membrane or soluble form of APRIL. In another preferred embodiment antibodies of the invention (including antibody fragments and variants) increase or enhance immunoglobulin production induced by the soluble form of BLyS. In another preferred embodiment antibodies of the invention (including antibody fragments and variants) increase or enhance immunoglobulin production induced by the membrane bound or soluble form of APRIL. In another preferred embodiment antibodies of the invention (including antibody fragments and variants) increase or enhance immunoglobulin production in response to T cell dependent immunogens. In another preferred embodiment antibodies of the invention (including antibody fragments and variants, and anti-antibody antibodies) increase or enhance immunoglobulin production in response to T cell independent immunogens.

[0407] In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate immune disorders. Immune disorders include, but are not limited to, autoimmune disorders (e.g., arthritis, graft rejection, Hashimoto's thyroiditis, insulin-dependent diabetes, lupus, idiopathic thrombocytopenic purpura, systemic lupus erythrematosus and multiple sclerosis), elective IgA deficiency, ataxia-telangiectasia, common variable immunodeficiency (CVID), X-linked agammaglobulinemia, severe combined immunodeficiency (SCID), Wiskott-Aldrich syndrome, idiopathic hyper-eosinophilic syndrome, monocytic leukemoid reaction, monocytic leukocytosis, monocytic leukopenia, monocytopenia, monocytosis, and graft or transplant rejection.

[0408] As discussed herein, antibodies and antibody compositions of the

invention, may be used to treat, prevent, ameliorate, diagnose or prognose various immune system-related disorders and/or conditions associated with these disorders, in mammals, preferably humans. Many autoimmune disorders result from inappropriate recognition of self as foreign material by immune cells. This inappropriate recognition results in an immune response leading to the destruction of the host tissue. Therefore, the administration of antibody and antibody compositions of the invention that can inhibit an immune response, particularly the proliferation of B cells and/or the production of immunoglobulins, may be an effective therapy in treating and/or preventing autoimmune disorders. Thus, in preferred embodiments, antibodies and antibody compositions of the invention are used to treat, prevent, ameliorate, diagnose and/or prognose an autoimmune disorder, or condition(s) associated with such disorder.

[0409] Autoimmune disorders and conditions associated with these disorders that may be treated, prevented, ameliorated, diagnosed and/or prognosed with the therapeutic and pharmaceutical compositions of the invention include, but are not limited to, autoimmune hemolytic anemia, autoimmune neonatal thrombocytopenia, idiopathic thrombocytopenia purpura, autoimmune neutropenia, autoimmunocytopenia, hemolytic anemia, antiphospholipid syndrome, dermatitis, gluten-sensitive enteropathy, allergic encephalomyelitis, myocarditis, relapsing polychondritis, rheumatic heart disease, glomerulonephritis (e.g., IgA nephropathy), Multiple Sclerosis, Neuritis, Uveitis Ophthalmia, Polyendocrinopathies, Purpura (e.g., Henloch-Scoenlein purpura), Reiter's Disease, Stiff-Man Syndrome, Autoimmune Pulmonary Inflammation, myocarditis, IgA glomerulonephritis, dense deposit disease, rheumatic heart disease, Guillain-Barre Syndrome, insulin dependent diabetes mellitis, and autoimmune inflammatory eye disease.

[0410] Additional autoimmune disorders and conditions associated with these disorders that may be treated, prevented, ameliorated, diagnosed and/or prognosed with the therapeutic and pharmaceutical compositions of the invention include, but are not limited to, autoimmune thyroiditis, hypothyroidism (i.e., Hashimoto's thyroiditis) (often characterized, e.g., by cell-mediated and humoral thyroid cytotoxicity), systemic lupus erhythematosus (often characterized, e.g., by circulating and locally generated immune complexes), discoid lupus, Goodpasture's syndrome (often characterized, e.g., by anti-basement membrane antibodies), Pemphigus (often characterized, e.g., by epidermal acantholytic antibodies), Receptor autoimmunities such as, for example, (a) Graves'

Disease (often characterized, e.g., by TSH receptor antibodies), (b) Myasthenia Gravis (often characterized, e.g., by acetylcholine receptor antibodies), and (c) insulin resistance (often characterized, e.g., by insulin receptor antibodies), autoimmune hemolytic anemia (often characterized, e.g., by phagocytosis of antibody-sensitized RBCs), autoimmune thrombocytopenic purpura (often characterized, e.g., by phagocytosis of antibody-sensitized platelets.

Additional autoimmune disorders and conditions associated with these [0411] disorders that may be treated, prevented, ameliorated, diagnosed and/or prognosed with the therapeutic and pharmaceutical compositions of the invention include, but are not limited to, rheumatoid arthritis (often characterized, e.g., by immune complexes in joints). schleroderma with anti-collagen antibodies (often characterized, e.g., by nucleolar and other nuclear antibodies), mixed connective tissue disease (often characterized, e.g., by antibodies to extractable nuclear antigens (e.g., ribonucleoprotein)), polymyositis/dermatomyositis (often characterized, e.g., by nonhistone ANA), pernicious anemia (often characterized, e.g., by antiparietal cell, microsomes, and intrinsic factor antibodies), idiopathic Addison's disease (often characterized, e.g., by humoral and cellmediated adrenal cytotoxicity, infertility (often characterized, e.g., by antispermatozoal antibodies), glomerulonephritis (often characterized, e.g., by glomerular basement membrane antibodies or immune complexes) such as primary glomerulonephritis and IgA nephropathy; bullous pemphigoid (often characterized, e.g., by IgG and complement in basement membrane), Sjögren's syndrome (often characterized, e.g., by multiple tissue antibodies, and/or a specific nonhistone ANA (SS-B)), diabetes millitus (often characterized, e.g., by cell-mediated and humoral islet cell antibodies), and adrenergic drug resistance (including adrenergic drug resistance with asthma or cystic fibrosis) (often characterized, e.g., by beta-adrenergic receptor antibodies), chronic active hepatitis (often characterized, e.g., by smooth muscle antibodies), primary biliary cirrhosis (often characterized, e.g., by mitchondrial antibodies), other endocrine gland failure (often characterized, e.g., by specific tissue antibodies in some cases), vitiligo (often characterized, e.g., by melanocyte antibodies), vasculitis (often characterized, e.g., by Ig and complement in vessel walls and/or low serum complement), post-MI (often characterized, e.g., by myocardial antibodies), cardiotomy syndrome (often characterized, e.g., by myocardial antibodies), urticaria (often characterized, e.g., by IgG and IgM

antibodies to IgE), atopic dermatitis (often characterized, e.g., by IgG and IgM antibodies to IgE), asthma (often characterized, e.g., by IgG and IgM antibodies to IgE), inflammatory myopathies, and many other inflammatory, granulomatous, degenerative, and atrophic disorders.

[0412] In a preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, a member of the group: autoimmune hemolytic anemia, as primary glomerulonephritis, IgA glomerulonephritis, Goodpasture's syndrome, idiopathic thrombocytopenia, Multiple Sclerosis, Myasthenia Gravis, Pemphigus, polymyositis/dermatomyositis, relapsing polychondritis, rheumatoid arthritis, Sjögren's syndrome, systemic lupus erythematosus, Uveitis, vasculitis, and primary biliary cirrhosis.

[0413] In another preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, an immune based-rheumatologic disease, such as, for example, SLE, rheumatoid arthritis, CREST syndrome (a variant of scleroderma characterized by calcinosis, Raynaud's phenomenon, esophageal motility disorders, sclerodactyly, and telangiectasia.), Seronegative spondyloarthropathy (SpA), polymyositis/ dermatomyositis, microscopic polyangiitis, hepatitis C-associated arthritis, Takayasu's arteritis, and undifferentiated connective tissue disorder.

[0414] In a specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, rheumatoid arthritis and/or medical conditions associated therewith.

[0415] For example, an antibody, or antibodies, of the present invention are used to treat patients with clinical diagnosis of rheumatoid arthritis (RA). The patient treated preferably will not have a B cell malignancy. Moreover, the patient is optionally further treated with any one or more agents employed for treating RA such as salicylate; nonsteroidal anti-inflammatory drugs such as indomethacin, phenylbutazone, phenylacetic acid derivatives (e.g. ibuprofen and fenoprofen), naphthalene acetic acids (naproxen), pyrrolealkanoic acid (tometin), indoleacetic acids (sulindac), halogenated anthranilic acid (meclofenamate sodium), piroxicam, zomepirac and diflunisal; antimalarials such as

chloroquine; gold salts; penicillamine; or immunosuppressive agents such as methotrexate or corticosteroids in dosages known for such drugs or reduced dosages. Preferably however, the patient is only treated with an antibody, or antibodies, of the present invention. Antibodies of the present invention are administered to the RA patient according to a dosing schedule as described infra, which may be readily determined by one of ordinary skill in the art. The primary response is determined by the Paulus index (Paulus et al. Athritis Rheum. 33:477-484 (1990)), i.e. improvement in morning stiffness. number of painful and inflamed joints, erythrocyte sedimentation (ESR), and at least a 2-point improvement on a 5-point scale of disease severity assessed by patient and by physician. Administration of an antibody, or antibodies, of the present invention will alleviate one or more of the symptoms of RA in the patient treated as described above. [0416] In a specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, amelioate, diagnose or prognose, lupus and/or medical conditions associated therewith. Lupus-associated conditions that may be treated, prevented, ameliorated, prognosed and/or diagnosed with the antibodies and antibody compositions of the invention include, but are not limited to, hematologic disorders (e.g., hemolytic anemia, leukopenia, lymphopenia, and thrombocytopenia), immunologic disorders (e.g., anti-DNA antibodies, and anti-Sm antibodies), rashes,

and antibody compositions of the invention include, but are not limited to, hematologic disorders (e.g., hemolytic anemia, leukopenia, lymphopenia, and thrombocytopenia), immunologic disorders (e.g., anti-DNA antibodies, and anti-Sm antibodies), rashes, photosensitivity, oral ulcers, arthritis, fever, fatigue, weight loss, serositis (e.g., pleuritus (pleurisy)), renal disorders (e.g., nephritis), neurological disorders (e.g., seizures, peripheral neuropathy, CNS related disorders), gastroinstestinal disorders, Raynaud phenomenon, and pericarditis. In a preferred embodiment, therapeutic and pharmaceutical compositions of the invention are used to treat, prevent, ameliorate, diagnose, or prognose, renal disorders associated with systemic lupus erythematosus. In a most preferred embodiment, therapeutic and pharmaceutical compositions of the invention are used to treat, prevent, ameliorate, diagnose, or prognose, nephritis associated with systemic lupus erythematosus. In another most preferred embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate lupus or glomerular nephritis.

[0417] In a further specific embodiment, antibodies of the invention are used to treat, inhibit, prognose, diagnose or prevent hemolytic anemia. For example, patients diagnosed with autoimmune hemolytic anemia (AIHA), e.g., cryoglobinemia or Coombs positive anemia, are treated with an antibody, or antibodies, of the present invention. AIHA is an acquired hemolytic anemia due to auto-antibodies that react with the patient's red blood cells. The patient treated preferably will not have a B cell malignancy. Further adjunct therapies (such as glucocorticoids, prednisone, azathioprine, cyclophosphamide, vinca-laden platelets or Danazol) may be combined with the antibody therapy, but preferably the patient is treated with an antibody, or antibodies, of the present invention as a single-agent throughout the course of therapy. Antibodies of the present invention are administered to the hemolytic anemia patient according to a dosing schedule as described infra, which may be readily determined by one of ordinary skill in the art. Overall response rate is determined based upon an improvement in blood counts, decreased requirement for transfusions, improved hemoglobin levels and/or a decrease in the evidence of hemolysis as determined by standard chemical parameters. Administration of an antibody, or antibodies of the present invention will improve any one or more of the symptoms of hemolytic anemia in the patient treated as described above. For example, the patient treated as described above will show an increase in hemoglobin and an improvement in chemical parameters of hemolysis or return to normal as measured by serum lactic dehydrogenase and/or bilirubin.

[0418] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, Sjögren's Syndrome and/or medical conditions associated therewith.

[0419] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, HIV infection and/or medical conditions associated therewith (e.g. AIDS).

[0420] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, Myasthenia gravis and/or medical conditions associated therewith.

- [0421] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, IgA nephropathy and/or medical conditions associated therewith.
- [0422] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, hemolytic anemia and/or medical conditions associated therewith.
- [0423] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, thyroiditis and/or medical conditions associated therewith.
- [0424] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, Goodpasture's Syndrome and/or medical conditions associated therewith.
- [0425] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, multiple sclerosis and/or medical conditions associated therewith.
- [0426] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, chronic lymphocytic leukemia (CLL) and/or medical conditions associated therewith.
- [0427] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, multiple myeloma and/or medical conditions associated therewith.
- [0428] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, Non-Hodgkin's lymphoma and/or medical conditions associated therewith.
- [0429] In another specific preferred embodiment, therapeutic and pharmaceutical compositions of the invention, are used to treat, prevent, ameliorate, diagnose or prognose, Hodgkin's disease and/or medical conditions associated therewith.
- [0430] In another specific embodiment, antibodies of the invention are used to treat, inhibit, prognose, diagnose or prevent adult immune thrombocytopenic purpura.

Adult immune thrombocytopenic purpura (ITP) is a relatively rare hematologic disorder that constitutes the most common of the immune-mediated cytopenias. The disease typically presents with severe thrombocytopenia that may be associated with acute hemorrhage in the presence of normal to increased megakaryocytes in the bone marrow. Most patients with ITP have an IgG antibody directed against target antigens on the outer surface of the platelet membrane, resulting in platelet sequestration in the spleen and accelerated reticuloendothelial destruction of platelets (Bussell, J.B. Hematol. Oncol. Clin. North Am. (4):179 (1990)). A number of therapeutic interventions have been shown to be effective in the treatment of ITP. Steroids are generally considered first-line therapy, after which most patients are candidates for intravenous immunoglobulin (IVIG), splenectomy, or other medical therapies including vincristine or immunosuppressive/cytotoxic agents. Up to 80% of patients with ITP initially respond to a course of steroids, but far fewer have complete and lasting remissions. Splenectomy has been recommended as standard second-line therapy for steroid failures, and leads to prolonged remission in nearly 60% of cases yet may result in reduced immunity to infection. Splenectomy is a major surgical procedure that may be associated with substantial morbidity (15%) and mortality (2%). IVIG has also been used as second line medical therapy, although only a small proportion of adult patients with ITP achieve remission. Therapeutic options that would interfere with the production of autoantibodies by activated B cells without the associated morbidities that occur with corticosteroids and/or splenectomy would provide an important treatment approach for a proportion of patients with ITP. Patients with clinical diagnosis of ITP are treated with an antibody, or antibodies of the present invention, optionally in combination with steroid therapy. The patient treated will not have a B cell malignancy. Antibodies of the present invention are administered to the RA patient according to a dosing schedule as described infra, which may be readily determined by one of ordinary skill in the art. Overall patient response rate is determined based upon a platelet count determined on two consecutive occasions two weeks apart following

treatments as described above. See, George et al. "Idiopathic Thrombocytopenic Purpura:

A Practice Guideline Developed by Explicit Methods for The American Society of

Hematology", Blood 88:3-40 (1996), expressly incorporated herein by reference.

In another embodiment, therapeutic or pharmaceutical compositions of the [0431] invention are administered to an animal to treat, prevent or ameliorate an IgE-mediated allergic reaction or histamine-mediated allergic reaction. Examples of allergic reactions include, but are not limited to, asthma, rhinitis, eczema, chronic urticaria, and atopic dermatitis. In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent, or ameliorate anaphylaxis, hypersensitivity to an antigenic molecule, or blood group incompatibility. In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate or modulate inflammation or an inflammatory disorder. Examples of chronic and acute inflammatory disorders that may be treated prevented or ameliorated with the therapeutic and pharmaceutical compositions of the invention include, but are not limited to, chronic prostatitis, granulomatous prostatitis and malacoplakia, inflammation associated with infection (e.g., septic shock, sepsis, or systemic inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine induced lung injury, Crohn's disease, inflammatory bowel disease, chronic and acute inflammatory pulmonary diseases, bacterial infection, psoriasis, septicemia, cerebral malaria, arthritis, gastroenteritis, and glomerular nephritis.

[0432] In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate ischemia and arteriosclerosis. Examples of such disorders include, but are not limited to, reperfusion damage (e.g., in the heart and/or brain) and cardiac hypertrophy.

[0433] Therapeutic or pharmaceutical compositions of the invention, may also be administered to modulate blood clotting and to treat or prevent blood clotting disorders, such as, for example, antibody-mediated thrombosis (i.e., antiphospholipid antibody syndrome (APS)). For example, therapeutic or pharmaceutical compositions of the invention, may inhibit the proliferation and differentiation of cells involved in producing anticardiolipin antibodies. These compositions of the invention can be used to treat, prevent, ameliorate, diagnose, and/or prognose thrombotic related events including, but

not limited to, stroke (and recurrent stroke), heart attack, deep vein thrombosis, pulmonary embolism, myocardial infarction, coronary artery disease (e.g., antibody –mediated coronary artery disease), thrombosis, graft reocclusion following cardiovascular surgery (e.g., coronary arterial bypass grafts, recurrent fetal loss, and recurrent cardiovascular thromboembolic events.

Therapeutic or pharmaceutical compositions of the invention, may also be administered to treat, prevent, or ameliorate organ rejection or graft-versus-host disease (GVHD) and/or conditions associated therewith. Organ rejection occurs by host immune cell destruction of the transplanted tissue through an immune response. Similarly, an immune response is also involved in GVHD, but, in this case, the foreign transplanted immune cells destroy the host tissues. The administration of antibodies of the invention, that inhibit an immune response, may be an effective therapy in preventing organ rejection or GVHD.

In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate a disease or disorder diseases associated with increased apoptosis including, but not limited to, AIDS, neurodegenerative disorders (such as Alzheimer's disease, Parkinson's disease, Amyotrophic lateral sclerosis, Retinitis pigmentosa, Cerebellar degeneration), myelodysplastic syndromes (such as aplastic anemia), ischemic injury (such as that caused by myocardial infarction, stroke and reperfusion injury), toxin-induced liver disease (such as that caused by alcohol), septic shock, cachexia and anorexia. In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate bone marrow failure, for example, aplastic anemia and myelodysplastic syndrome.

In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate growth, progression, and/or metastases of malignancies and proliferative disorders associated with increased cell survival, or the inhibition of apoptosis. Examples of such disorders, include, but are not limited to, leukemia (e.g., acute leukemia such as acute lymphocytic leukemia and acute myelocytic leukemia), neoplasms, tumors (e.g., fibrosarcoma, myxosarcoma, liposarcoma, chondrosarcoma, osteogenic sarcoma, chordoma, angiosarcoma, endotheliosarcoma, lymphangiosarcoma, lymphangioendotheliosarcoma,

synovioma, mesothelioma, Ewing's tumor, leiomyosarcoma, rhabdomyosarcoma, colon carcinoma, pancreatic cancer, breast cancer, ovarian cancer, prostate cancer, squamous cell carcinoma, basal cell carcinoma, adenocarcinoma, sweat gland carcinoma, sebaceous gland carcinoma, papillary carcinoma, papillary adenocarcinomas, cystadenocarcinoma, medullary carcinoma, bronchogenic carcinoma, renal cell carcinoma, hepatoma, bile duct carcinoma, choriocarcinoma, seminoma, embryonal carcinoma, Wilms' tumor, cervical cancer, testicular tumor, lung carcinoma, small cell lung carcinoma, bladder carcinoma, epithelial carcinoma, glioma, astrocytoma, medulloblastoma, craniopharyngioma, ependymoma, pinealoma, hemangioblastoma, acoustic neuroma, oligodendroglioma, meningioma, melanoma, neuroblastoma, and retinoblastoma), heavy chain disease, metastases, or any disease or disorder characterized by uncontrolled cell growth.

[0437] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used to treat or prevent a disorder characterized by hpergammagloulinemia (e.g., AIDS, autoimmune diseases, and some immunodeficiencies).

In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used to treat or prevent a disorder characterized by deficient serum immunoglobulin production, recurrent infections, and/or immune system dysfunction. Moreover, therapeutic or pharmaceutical compositions of the invention may be used to treat or prevent infections of the joints, bones, skin, and/or parotid glands, blood-borne infections (e.g., sepsis, meningitis, septic arthritis, and/or osteomyelitis), autoimmune diseases (e.g., those disclosed herein), inflammatory disorders, and malignancies, and/or any disease or disorder or condition associated with these infections, diseases, disorders and/or malignancies) including, but not limited to, CVID, other primary immune deficiencies, HIV disease, CLL, recurrent bronchitis, sinusitis, otitis media, conjunctivitis, pneumonia, hepatitis, meningitis, herpes zoster (e.g., severe herpes zoster), and/or pneumocystis carnii.

[0439] Therapeutic or pharmaceutical compositions of the invention of the invention thereof, may be used to diagnose, prognose, treat or prevent one or more of the following diseases or disorders, or conditions associated therewith: primary immuodeficiencies, immune-mediated thrombocytopenia, Kawasaki syndrome, bone marrow transplant (e.g., recent bone marrow transplant in adults or children), chronic B-

cell lymphocytic leukemia, HIV infection (e.g., adult or pediatric HIV infection), chronic inflammatory demyelinating polyneuropathy, and post-transfusion purpura.

[0440] Additionally, therapeutic or pharmaceutical compositions of the invention may be used to diagnose, prognose, treat or prevent one or more of the following diseases, disorders, or conditions associated therewith, Guillain-Barre syndrome, anemia (e.g., anemia associated with parvovirus B19, patients with stable multiple myeloma who are at high risk for infection (e.g., recurrent infection), autoimmune hemolytic anemia (e.g., warm-type autoimmune hemolytic anemia), thrombocytopenia (e.g., neonatal thrombocytopenia), and immune-mediated neutropenia), transplantation (e.g., cytomegalovirus (CMV)-negative recipients of CMV-positive organs), hypogammaglobulinemia (e.g., hypogammaglobulinemic neonates with risk factor for infection or morbidity), epilepsy (e.g., intractable epilepsy), systemic vasculitic syndromes, myasthenia gravis (e.g., decompensation in myasthenia gravis), dermatomyositis, and polymyositis.

[0441] Additional preferred embodiments of the invention include, but are not limited to, the use of therapeutic or pharmaceutical compositions of the invention in the following applications:

[0442] Administration to an animal (e.g., mouse, rat, rabbit, hamster, guinea pig, pigs, micro-pig, chicken, camel, goat, horse, cow, sheep, dog, cat, non-human primate, and human, most preferably human) to boost the immune system to produce increased quantities of one or more antibodies (e.g., IgG, IgA, IgM, and IgE), to induce higher affinity antibody production (e.g., IgG, IgA, IgM, and IgE), and/or to increase an immune response. In a specific nonexclusive embodiment, therapeutic or pharmaceutical compositions of the invention are administered to boost the immune system to produce increased quantities of IgG. In another specific nonexclusive embodiment, antibodies of the are administered to boost the immune system to produce increased quantities of IgA. In another specific nonexclusive embodiment antibodies of the invention are administered to boost the immune system to produce increased quantities of IgM.

[0443] Administration to an animal (including, but not limited to, those listed above, and also including transgenic animals) incapable of producing functional endogenous antibody molecules or having an otherwise compromised endogenous immune system, but which is capable of producing human immunoglobulin molecules by

means of a reconstituted or partially reconstituted immune system from another animal (see, e.g., published PCT Application Nos. WO98/24893, WO/9634096, WO/9633735, and WO/9110741).

In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a vaccine adjuvant that enhances immune responsiveness to specific antigen. In a specific embodiment, the vaccine is an antibody described herein. In another specific embodiment, the vaccine adjuvant is a polynucleotide described herein (e.g., an antibody polynucleotide genetic vaccine adjuvant). As discussed herein, therapeutic or pharmaceutical compositions of the invention may be administered using techniques known in the art, including but not limited to, liposomal delivery, recombinant vector delivery, injection of naked DNA, and gene gun delivery.

[0445] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an adjuvant to enhance tumor-specific immune responses.

[0446] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an adjuvant to enhance anti-viral immune responses. Anti-viral immune responses that may be enhanced using the compositions of the invention as an adjuvant, include, but are not limited to, virus and virus associated diseases or symptoms described herein or otherwise known in the art. In specific embodiments, the compositions of the invention are used as an adjuvant to enhance an immune response to a virus, disease, or symptom selected from the group consisting of: AIDS, meningitis, Dengue, EBV, and hepatitis (e.g., hepatitis B). In another specific embodiment, the compositions of the invention are used as an adjuvant to enhance an immune response to a virus, disease, or symptom selected from the group consisting of: HIV/AIDS, Respiratory syncytial virus, Dengue, Rotavirus, Japanese B encephalitis, Influenza A and B. Parainfluenza, Measles, Cytomegalovirus, Rabies, Junin, Chikungunya, Rift Valley fever, Herpes simplex, and yellow fever. In another specific embodiment, the compositions of the invention are used as an adjuvant to enhance an immune response to the HIV gp120 antigen.

[0447] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an adjuvant to enhance anti-bacterial or anti-fungal immune responses. Anti-bacterial or anti-fungal immune responses that may be enhanced using the

compositions of the invention as an adjuvant, include bacteria or fungus and bacteria or fungus associated diseases or symptoms described herein or otherwise known in the art. In specific embodiments, the compositions of the invention are used as an adjuvant to enhance an immune response to a bacteria or fungus, disease, or symptom selected from the group consisting of: tetanus, Diphtheria, botulism, and meningitis type B. In another specific embodiment, the compositions of the invention are used as an adjuvant to enhance an immune response to a bacteria or fungus, disease, or symptom selected from the group consisting of: Vibrio cholerae, Mycobacterium leprae, Salmonella typhi, Salmonella paratyphi, Neisseria meningitidis, Streptococcus pneumoniae, Group B streptococcus, Shigella spp., Enterotoxigenic Escherichia coli, Enterohemorrhagic E. coli, Borrelia burgdorferi, and Plasmodium (malaria).

[0448] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an adjuvant to enhance anti-parasitic immune responses. Anti-parasitic immune responses that may be enhanced using the compositions of the invention as an adjuvant, include parasite and parasite associated diseases or symptoms described herein or otherwise known in the art. In specific embodiments, the compositions of the invention are used as an adjuvant to enhance an immune response to a parasite. In another specific embodiment, the compositions of the invention are used as an adjuvant to enhance an immune response to Plasmodium (malaria).

[0449] In a specific embodiment, compositions of the invention may be administered to patients as vaccine adjuvants. In a further specific embodiment, compositions of the invention may be administered as vaccine adjuvants to patients suffering from an immune-deficiency. In a further specific embodiment, compositions of the invention may be administered as vaccine adjuvants to patients suffering from HIV.

[0450] In a specific embodiment, compositions of the invention may be used to increase or enhance antigen-specific antibody responses to standard and experimental vaccines. In a specific embodiment, compositions of the invention may be used to enhance seroconversion in patients treated with standard and experimental vaccines. In another specific embodiment, compositions of the invention may be used to increase the repertoire of antibodies recognizing unique epitopes in response to standard and experimental vaccination.

[0451] In a preferred embodiment, antibodies of the invention (including antibody

fragments and variants, and anti-antibody antibodies) increase or enhance antigen-specific antibody responses to standard and experimental vaccines by regulating binding of the soluble form of BLyS to a BLyS receptor (e.g., BCMA and TACI). In another preferred embodiment, antibodies of the invention (including antibody fragments and variants, and anti-antibody antibodies) increase or enhance antigen-specific antibody responses to standard and experimental vaccines by regulating binding of the soluble form of APRIL to an APRIL receptor (e.g., BCMA and TACI).

In a preferred embodiment, antibodies of the invention (including antibody fragments and variants, and anti-antibody antibodies) increase or enhance seroconversion in patients treated with standard and experimental vaccines by regulating binding of the soluble form of BLyS to BLyS receptor (e.g., BCMA and TACI). In another preferred embodiment, antibodies of the invention (including antibody fragments and variants, and anti-antibody antibodies) increase or enhance seroconversion in patients treated with standard and experimental vaccines by regulating binding of the soluble form of APRIL to an APRIL receptor (e.g., BCMA and TACI).

[0453] In a preferred embodiment, antibodies of the invention (including antibody fragments and variants, and anti-antibody antibodies) increase or enhance the repertoire of antibodies recognizing unique epitopes in response to standard and experimental vaccination by regulating binding of the soluble form of BLyS to a BLyS receptor (e.g., BCMA and TACI). In another preferred embodiment, antibodies of the invention (including antibody fragments and variants, and anti-antibody antibodies) increase or enhance the repertoire of antibodies recognizing unique epitopes in response to standard and experimental vaccination by regulating binding of the soluble form of APRIL to an APRIL receptor (e.g., BCMA and TACI).

[0454] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a stimulator of B cell responsiveness to pathogens.

[0455] In a specific embodiment, therapeutic or pharmaceutical compositions of

the invention are used as an agent that elevates the immune status of an individual prior to their receipt of immunosuppressive therapies.

[0456] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an agent to induce higher affinity antibodies.

[0457] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an agent to increase serum immunoglobulin concentrations.

[0458] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an agent to accelerate recovery of immunocompromised individuals.

[0459] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an agent to boost immunoresponsiveness among aged populations.

[0460] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an immune system enhancer prior to, during, or after bone marrow transplant and/or other transplants (e.g., allogeneic or xenogeneic organ transplantation). With respect to transplantation, compositions of the invention may be administered prior to, concomitant with, and/or after transplantation. In a specific embodiment, compositions of the invention are administered after transplantation, prior to the beginning of recovery of T-cell populations. In another specific embodiment, compositions of the invention are first administered after transplantation after the beginning of recovery of T cell populations, but prior to full recovery of B cell populations.

In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an agent to boost immunoresponsiveness among B cell immunodeficient individuals, such as, for example, an individual who has undergone a partial or complete splenectomy. B cell immunodeficiencies that may be ameliorated or treated by administering the antibodies and/or compositions of the invention include, but are not limited to, severe combined immunodeficiency (SCID)-X linked, SCID-autosomal, adenosine deaminase deficiency (ADA deficiency), X-linked agammaglobulinemia (XLA), Bruton's disease, congenital agammaglobulinemia, X-linked infantile

agammaglobulinemia, acquired agammaglobulinemia, adult onset agammaglobulinemia, late-onset agammaglobulinemia, dysgammaglobulinemia, hypogammaglobulinemia, transient hypogammaglobulinemia of infancy, unspecified hypogammaglobulinemia, agammaglobulinemia, common variable immunodeficiency (CVID) (acquired), Wiskott-Aldrich Syndrome (WAS), X-linked immunodeficiency with hyper IgM, non X-linked immunodeficiency with hyper IgM, selective IgA deficiency, IgG subclass deficiency (with or without IgA deficiency), antibody deficiency with normal or elevated Igs, immunodeficiency with thymoma, Ig heavy chain deletions, kappa chain deficiency, B cell lymphoproliferative disorder (BLPD), selective IgM immunodeficiency, recessive agammaglobulinemia (Swiss type), reticular dysgenesis, neonatal neutropenia, severe congenital leukopenia, thymic alymphoplasia-aplasia or dysplasia with immunodeficiency, ataxia-telangiectasia, short limbed dwarfism, X-linked lymphoproliferative syndrome (XLP), Nezelof syndrome-combined immunodeficiency with Igs, purine nucleoside phosphorylase deficiency (PNP), MHC Class II deficiency (Bare Lymphocyte Syndrome) and severe combined immunodeficiency.

[0462] In a specific embodiment, antibodies and/or compositions of the invention are administered to treat or ameliorate selective IgA deficiency.

[0463] In another specific embodiment, antibodies and/or compositions of the invention are administered to treat or ameliorate ataxia-telangiectasia.

[0464] In another specific embodiment antibodies and/or compositions of the invention are administered to treat or ameliorate common variable immunodeficiency.

[0465] In another specific embodiment, antibodies and/or compositions of the invention are administered to treat or ameliorate X-linked agammaglobulinemia.

[0466] In another specific embodiment, antibodies and/or compositions of the invention are administered to treat or ameliorate severe combined immunodeficiency (SCID).

[0467] In another specific embodiment, antibodies and/or compositions of the invention are administered to treat or ameliorate Wiskott-Aldrich syndrome.

[0468] In another specific embodiment, antibodies and/or compositions of the invention are administered to treat or ameliorate X-linked Ig deficiency with hyper IgM.

[0469] As an agent to boost immunoresponsiveness among individuals having an acquired loss of B cell function. Conditions resulting in an acquired loss of B cell

function that may be ameliorated or treated by administering antibodies and/or compositions of the invention include, but are not limited to, HIV Infection, AIDS, bone marrow transplant, and B cell chronic lymphocytic leukemia (CLL).

[0470] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an agent to boost immunoresponsiveness among individuals having a temporary immune deficiency. Conditions resulting in a temporary immune deficiency that may be ameliorated or treated by administering antibodies and/or compositions of the invention include, but are not limited to, recovery from viral infections (e.g., influenza), conditions associated with malnutrition, recovery from infectious mononucleosis, or conditions associated with stress, recovery from measles, recovery from blood transfusion, recovery from surgery.

[0471] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a regulator of antigen presentation by monocytes, dendritic cells, T cells and/or B-cells. In one embodiment, antibody polypeptides or polynucleotides enhance antigen presentation or antagonize antigen presentation in vitro or in vivo. Moreover, in related embodiments, this enhancement or antagonization of antigen presentation may be useful in anti-tumor treatment or to modulate the immune system.

[0472] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a mediator of mucosal immune responses. The expression of BLyS on monocytes, the expression of BLyS receptor on B cells, and the responsiveness of B cells to BLyS suggests that it may be involved in exchange of signals between B cells and monocytes or their differentiated progeny. This activity is in many ways analogous to the CD40-CD154 signalling between B cells and T cells. Anti-BLyS antibodies and compositions of the invention may therefore be good regulators of T cell independent immune responses to environmental pathogens. In particular, the unconventional B cell populations (CD5+) that are associated with mucosal sites and responsible for much of the innate immunity in humans may respond to antibodies or compositions of the invention thereby enhancing or inhibiting individual's immune status.

[0473] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an agent to direct an individual's immune system towards development of a humoral response (i.e. TH2) as opposed to a TH1 cellular response.

[0474] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a means to induce tumor proliferation and thus make it more susceptible to anti-neoplastic agents. For example, multiple myeloma is a slowly dividing disease and is thus refractory to virtually all anti-neoplastic regimens. If these cells were forced to proliferate more rapidly, their susceptibility profile would likely change.

[0475] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a monocyte cell specific binding protein to which specific activators or inhibitors of cell growth may be attached. The result would be to focus the activity of such activators or inhibitors onto normal, diseased, or neoplastic B cell populations.

[0476] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a B cell specific binding protein to which specific activators or inhibitors of cell growth may be attached. The result would be to focus the activity of such activators or inhibitors onto normal, diseased, or neoplastic B cell populations.

[0477] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a means of detecting monocytic cells by virtue of its specificity. This application may require labeling the protein with biotin or other agents (e.g., as described herein) to afford a means of detection.

[0478] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a means of detecting B-lineage cells by virtue of its specificity. This application may require labeling the protein with biotin or other agents (e.g., as described herein) to afford a means of detection.

[0479] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a stimulator of B cell production in pathologies such as AIDS, chronic lymphocyte disorder and/or Common Variable immunodeficiency.

[0480] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as part of a monocyte selection device the function of which is to isolate monocytes from a heterogeneous mixture of cell types. Antibodies of the invention could be coupled to a solid support to which monocytes would then specifically bind.

Unbound cells would be washed out and the bound cells subsequently eluted. A nonlimiting use of this selection would be to allow purging of tumor cells from, for example, bone marrow or peripheral blood prior to transplant.

In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as part of a B cell selection device the function of which is to isolate B cells from a heterogeneous mixture of cell types. Antibodies of the invention (that do not inhibit BLyS/BLyS Receptor interaction) binding soluble BLyS could be coupled to a solid support to which B cells would then specifically bind. Unbound cells would be washed out and the bound cells subsequently eluted. A non-limiting use of this selection would be to allow purging of tumor cells from, for example, bone marrow or peripheral blood prior to transplant.

[0482] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a therapy for generation and/or regeneration of lymphoid tissues following surgery, trauma or genetic defect.

[0483] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a gene-based therapy for genetically inherited disorders resulting in immuno-incompetence such as observed among SCID patients.

[0484] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as an antigen for the generation of antibodies to inhibit or enhance BLyS mediated responses.

[0485] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a means of activating monocytes/macrophages to defend against parasitic diseases that effect monocytes such as Leishmania.

[0486] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as pretreatment of bone marrow samples prior to transplant. Such treatment would increase B cell representation and thus accelerate recovery.

[0487] In a specific embodiment, therapeutic or pharmaceutical compositions of the invention are used as a means of regulating secreted cytokines that are elicited by BLyS and/or BLyS receptor.

[0488] Antibody polypeptides or polynucleotides of the invention may be used to modulate IgE concentrations in vitro or in vivo.

[0489] Additionally, antibody polypeptides or polynucleotides of the invention may be used to treat, prevent, and/or diagnose IgE-mediated allergic reactions. Such allergic reactions include, but are not limited to, asthma, rhinitis, and eczema.

[0490] In a specific embodiment, antibody polypeptides or polynucleotides of the invention, are administered to treat, prevent, diagnose, and/or ameliorate selective IgA deficiency.

[0491] In another specific embodiment antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate ataxiatelangiectasia.

[0492] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate common variable immunodeficiency.

[0493] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate X-linked agammaglobulinemia.

[0494] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate severe combined immunodeficiency (SCID).

[0495] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate Wiskott-Aldrich syndrome.

[0496] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate X-linked Ig deficiency with hyper IgM. In a specific embodiment antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate X-linked Ig deficiency with hyper IgM.

[0497] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, and/or diagnose chronic myelogenous leukemia, acute myelogenous leukemia, leukemia, hystiocytic leukemia, monocytic leukemia (e.g., acute monocytic leukemia), leukemic reticulosis, Shilling Type monocytic

leukemia, and/or other leukemias derived from monocytes and/or monocytic cells and/or tissues.

[0498] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate monocytic leukemoid reaction, as seen, for example, with tuberculosis.

[0499] In another specific embodiment, antibody polypeptides or polynucleotides of the invention are administered to treat, prevent, diagnose, and/or ameliorate monocytic leukocytosis, monocytic leukopenia, monocytopenia, and/or monocytosis.

[0500] In a specific embodiment, antibody polypeptides or polynucleotides of the invention are used to treat, prevent, detect, and/or diagnose monocyte disorders and/or diseases, and/or conditions associated therewith.

In a specific embodiment, antibody polypeptides or polynucleotides of the invention are used to treat, prevent, detect, and/or diagnose primary B lymphocyte disorders and/or diseases, and/or conditions associated therewith. In one embodiment, such primary B lymphocyte disorders, diseases, and/or conditions are characterized by a complete or partial loss of humoral immunity. Primary B lymphocyte disorders, diseases, and/or conditions associated therewith that are characterized by a complete or partial loss of humoral immunity and that may be prevented, treated, detected and/or diagnosed with compositions of the invention include, but are not limited to, X-Linked Agammaglobulinemia (XLA), severe combined immunodeficiency disease (SCID), and selective IgA deficiency.

In a preferred embodiment antibody polypeptides or polynucleotides of the invention are used to treat, prevent, and/or diagnose diseases or disorders affecting or conditions associated with any one or more of the various mucous membranes of the body. Such diseases or disorders include, but are not limited to, for example, mucositis, mucoclasis, mucocolitis, mucocutaneous leishmaniasis (such as, for example, American leishmaniasis, leishmaniasis americana, nasopharyngeal leishmaniasis, and New World leishmaniasis), mucocutaneous lymph node syndrome (for example, Kawasaki disease), mucoenteritis, mucoepidermoid carcinoma, mucoepidermoid tumor, mucoepithelial dysplasia, mucoid adenocarcinoma, mucoid degeneration, myxoid degeneration; myxomatous degeneration; myxomatosis, mucoid medial degeneration (for example, cystic medial necrosis), mucolipidosis (including, for example, mucolipidosis I,

mucolipidosis II, mucolipidosis III, and mucolipidosis IV), mucolysis disorders, mucomembranous enteritis, mucoenteritis, mucopolysaccharidosis (such as, for example, type I mucopolysaccharidosis (i.e., Hurler's syndrome), type IS mucopolysaccharidosis (i.e., Scheie's syndrome or type V mucopolysaccharidosis), type II mucopolysaccharidosis (i.e., Hunter's syndrome), type III mucopolysaccharidosis (i.e., Sanfilippo's syndrome), type IV mucopolysaccharidosis (i.e., Morquio's syndrome), type VI mucopolysaccharidosis (i.e., Maroteaux-Lamy syndrome), type VII mucopolysaccharidosis (i.e., mucopolysaccharidosis due to beta-glucuronidase deficiency). and mucosulfatidosis), mucopolysacchariduria, mucopurulent conjunctivitis, mucopus, mucormycosis (i.e., zygomycosis), mucosal disease (i.e., bovine virus diarrhea), mucous colitis (such as, for example, mucocolitis and myxomembranous colitis), and mucoviscidosis (such as, for example, cystic fibrosis, cystic fibrosis of the pancreas, Clarke-Hadfield syndrome, fibrocystic disease of the pancreas, mucoviscidosis, and viscidosis). In a highly preferred embodiment, antibody polypeptides or polynucleotides of the invention are used to treat, prevent, and/or diagnose mucositis, especially as associated with chemotherapy.

[0503] In a preferred embodiment, antibody polypeptides or polynucleotides of the invention are used to treat, prevent, and/or diagnose diseases or disorders affecting or conditions associated with sinusitis.

[0504] An additional condition, disease or symptom that can be treated, prevented, and/or diagnosed by antibody polypeptides or polynucleotides of the invention is osteomyelitis.

[0505] An additional condition, disease or symptom that can be treated, prevented, and/or diagnosed by antibody polypeptides or polynucleotides of the invention is endocarditis.

[0506] All of the above described applications as they may apply to veterinary medicine.

[0507] Antibody polypeptides or polynucleotides of the invention may be used to treat, prevent, and/or diagnose diseases and disorders of the pulmonary system (e.g., bronchi such as, for example, sinopulmonary and bronchial infections and conditions associated with such diseases and disorders and other respiratory diseases and disorders. In specific embodiments, such diseases and disorders include, but are not limited to,

bronchial adenoma, bronchial asthma, pneumonia (such as, e.g., bronchial pneumonia, bronchopneumonia, and tuberculous bronchopneumonia), chronic obstructive pulmonary disease (COPD), bronchial polyps, bronchiectasia (such as, e.g., bronchiectasia sicca, cylindrical bronchiectasis, and saccular bronchiectasis), bronchiolar adenocarcinoma, bronchiolar carcinoma, bronchiolitis (such as, e.g., exudative bronchiolitis, bronchiolitis fibrosa obliterans, and proliferative bronchiolitis), bronchiolo-alveolar carcinoma, bronchitic asthma, bronchitis (such as, e.g., asthmatic bronchitis, Castellani's bronchitis, chronic bronchitis, croupous bronchitis, fibrinous bronchitis, hemorrhagic bronchitis, infectious avian bronchitis, obliterative bronchitis, plastic bronchitis, pseudomembranous bronchitis, putrid bronchitis, and verminous bronchitis), bronchocentric granulomatosis, bronchoedema, bronchoesophageal fistula, bronchogenic carcinoma, bronchogenic cyst, broncholithiasis, bronchomalacia, bronchomycosis (such as, e.g., bronchopulmonary aspergillosis), bronchopulmonary spirochetosis, hemorrhagic bronchitis, bronchorrhea, bronchospasm, bronchostaxis, bronchostenosis, Biot's respiration, bronchial respiration, Kussmaul respiration, Kussmaul-Kien respiration, respiratory acidosis, respiratory alkalosis, respiratory distress syndrome of the newborn, respiratory insufficiency, respiratory scleroma, respiratory syncytial virus, and the like.

[0508] In a specific embodiment, antibody polypeptides or polynucleotides of the invention are used to treat, prevent, and/or diagnose chronic obstructive pulmonary disease (COPD).

[0509] In another embodiment, antibody polypeptides or polynucleotides of the invention are used to treat, prevent, and/or diagnose fibroses and conditions associated with fibroses, including, but not limited to, cystic fibrosis (including such fibroses as cystic fibrosis of the pancreas, Clarke-Hadfield syndrome, fibrocystic disease of the pancreas, mucoviscidosis, and viscidosis), endomyocardial fibrosis, idiopathic retroperitoneal fibrosis, leptomeningeal fibrosis, mediastinal fibrosis, nodular subepidermal fibrosis, pericentral fibrosis, perimuscular fibrosis, pipestem fibrosis, replacement fibrosis, subadventitial fibrosis, and Symmers' clay pipestem fibrosis.

[0510] In another embodiment, therapeutic or pharmaceutical compositions of the invention are administered to an animal to treat, prevent or ameliorate infectious diseases. Infectious diseases include diseases associated with yeast, fungal, viral and bacterial infections. Viruses causing viral infections which can be treated or prevented in

accordance with this invention include, but are not limited to, retroviruses (e.g., human T-cell lymphotrophic virus (HTLV) types I and II and human immunodeficiency virus (HIV)), herpes viruses (e.g., herpes simplex virus (HSV) types I and II, Epstein-Barr virus, HHV6-HHV8, and cytomegalovirus), arenavirues (e.g., lassa fever virus), paramyxoviruses (e.g., morbillivirus virus, human respiratory syncytial virus, mumps, and pneumovirus), adenoviruses, bunyaviruses (e.g., hantavirus), cornaviruses, filoviruses (e.g., Ebola virus), flaviviruses (e.g., hepatitis C virus (HCV), yellow fever virus, and Japanese encephalitis virus), hepadnaviruses (e.g., hepatitis B viruses (HBV)). orthomyoviruses (e.g., influenza viruses A, B and C), papovaviruses (e.g., papillomavirues), picornaviruses (e.g., rhinoviruses, enteroviruses and hepatitis A viruses). poxviruses, reoviruses (e.g., rotaviruses), togaviruses (e.g., rubella virus), rhabdoviruses (e.g., rabies virus). Microbial pathogens causing bacterial infections include, but are not limited to, Streptococcus pyogenes, Streptococcus pneumoniae, Neisseria gonorrhoea, Neisseria meningitidis, Corynebacterium diphtheriae, Clostridium botulinum. Clostridium perfringens, Clostridium tetani, Haemophilus influenzae, Klebsiella pneumoniae, Klebsiella ozaenae, Klebsiella rhinoscleromotis, Staphylococcus aureus, Vibrio cholerae, Escherichia coli, Pseudomonas aeruginosa, Campylobacter (Vibrio) fetus, Campylobacter jejuni, Aeromonas hydrophila, Bacillus cereus, Edwardsiella tarda. Yersinia enterocolitica, Yersinia pestis, Yersinia pseudotuberculosis, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Salmonella typhimurium, Treponema pallidum, Treponema pertenue, Treponema carateneum, Borrelia vincentii, Borrelia burgdorferi, Leptospira icterohemorrhagiae, Mycobacterium tuberculosis, Toxoplasma gondii, Pneumocystis carinii, Francisella tularensis, Brucella abortus, Brucella suis, Brucella melitensis, Mycoplasma spp., Rickettsia prowazeki, Rickettsia tsutsugumushi, Chlamydia spp., and Helicobacter pylori.

Gene Therapy

[0511] In a specific embodiment, nucleic acids comprising sequences encoding antibodies or functional derivatives thereof, are administered to treat, inhibit or prevent a disease or disorder associated with aberrant expression and/or activity of BLyS and/or its receptor, by way of gene therapy. Gene therapy refers to therapy performed by the

administration to a subject of an expressed or expressible nucleic acid. In this embodiment of the invention, the nucleic acids produce their encoded protein that mediates a therapeutic effect.

[0512] Any of the methods for gene therapy available in the art can be used according to the present invention. Exemplary methods are described below.

[0513] For general reviews of the methods of gene therapy, see Goldspiel et al., Clinical Pharmacy 12:488-505 (1993); Wu and Wu, Biotherapy 3:87-95 (1991); Tolstoshev, Ann. Rev. Pharmacol. Toxicol. 32:573-596 (1993); Mulligan, Science 260:926-932 (1993); and Morgan and Anderson, Ann. Rev. Biochem. 62:191-217 (1993); May, TIBTECH 1 1(5):155-215 (1993). Methods commonly known in the art of recombinant DNA technology which can be used are described in Ausubel et al. (eds.), Current Protocols in Molecular Biology, John Wiley & Sons, NY (1993); and Kriegler, Gene Transfer and Expression, A Laboratory Manual, Stockton Press, NY (1990).

alternatively consists of, nucleic acids encoding an antibody, said nucleic acids being part of an expression vector that expresses the antibody or fragments or chimeric proteins or heavy or light chains thereof in a suitable host. In particular, such nucleic acids have promoters, preferably heterologous promoters, operably linked to the antibody coding region, said promoter being inducible or constitutive, and, optionally, tissue-specific. In another particular embodiment, nucleic acid molecules are used in which the antibody coding sequences and any other desired sequences are flanked by regions that promote homologous recombination at a desired site in the genome, thus providing for intrachromosomal expression of the antibody encoding nucleic acids (Koller and Smithies, Proc. Natl. Acad. Sci. USA 86:8932-8935 (1989); Zijlstra et al., Nature 342:435-438 (1989). In specific embodiments, the expressed antibody molecule is an scFv; alternatively, the nucleic acid sequences include sequences encoding both the heavy and light chains, or fragments or variants thereof, of an antibody.

[0515] Delivery of the nucleic acids into a patient may be either direct, in which case the patient is directly exposed to the nucleic acid or nucleic acid- carrying vectors, or indirect, in which case, cells are first transformed with the nucleic acids in vitro, then transplanted into the patient. These two approaches are known, respectively, as in vivo or ex vivo gene therapy.

[0516] In a specific embodiment, the nucleic acid sequences are directly administered in vivo, where it is expressed to produce the encoded product. This can be accomplished by any of numerous methods known in the art, e.g., by constructing them as part of an appropriate nucleic acid expression vector and administering it so that they become intracellular, e.g., by infection using defective or attenuated retrovirals or other viral vectors (see U.S. Patent No. 4,980,286), or by direct injection of naked DNA, or by use of microparticle bombardment (e.g., a gene gun; Biolistic, Dupont), or coating with lipids or cell-surface receptors or transfecting agents, encapsulation in liposomes, microparticles, or microcapsules, or by administering them in linkage to a peptide which is known to enter the nucleus, by administering it in linkage to a ligand subject to receptormediated endocytosis (see, e.g., Wu and Wu, J. Biol. Chem. 262:4429-4432 (1987)) (which can be used to target cell types specifically expressing the receptors), etc. In another embodiment, nucleic acid-ligand complexes can be formed in which the ligand comprises a fusogenic viral peptide to disrupt endosomes, allowing the nucleic acid to avoid lysosomal degradation. In yet another embodiment, the nucleic acid can be targeted in vivo for cell specific uptake and expression, by targeting a specific receptor (see, e.g., PCT Publications WO 92/06 180; WO 92/22635; W092/203 16; W093/14188, WO 93/20221). Alternatively, the nucleic acid can be introduced intracellularly and incorporated within host cell DNA for expression, by homologous recombination (Koller and Smithies, Proc. Natl. Acad. Sci. USA 86:8932-8935 (1989); Zijlstra et al., Nature 342:435-438 (1989)).

[0517] In a specific embodiment, viral vectors that contains nucleic acid sequences encoding an antibody of the invention or fragments or variants thereof are used. For example, a retroviral vector can be used (see Miller et al., Meth. Enzymol. 217:581-599 (1993)). These retroviral vectors contain the components necessary for the correct packaging of the viral genome and integration into the host cell DNA. The nucleic acid sequences encoding the antibody to be used in gene therapy are cloned into one or more vectors, which facilitates delivery of the gene into a patient. More detail about retroviral vectors can be found in Boesen et al., Biotherapy 6:29 1-302 (1994), which describes the use of a retroviral vector to deliver the mdr 1 gene to hematopoietic stem cells in order to make the stem cells more resistant to chemotherapy. Other references illustrating the use of retroviral vectors in gene therapy are: Clowes et al., J. Clin. Invest. 93:644-651(1994);

Klein et al., Blood 83:1467-1473 (1994); Salmons and Gunzberg, Human Gene Therapy 4:129-141 (1993); and Grossman and Wilson, Curr. Opin. in Genetics and Devel. 3:110-114 (1993).

Adenoviruses are other viral vectors that can be used in gene therapy. Adenoviruses are especially attractive vehicles for delivering genes to respiratory epithelia. Adenoviruses naturally infect respiratory epithelia where they cause a mild disease. Other targets for adenovirus-based delivery systems are liver, the central nervous system, endothelial cells, and muscle. Adenoviruses have the advantage of being capable of infecting non-dividing cells. Kozarsky and Wilson, Current Opinion in Genetics and Development 3:499-503 (1993) present a review of adenovirus-based gene therapy. Bout et al., Human Gene Therapy 5:3-10 (1994) demonstrated the use of adenovirus vectors to transfer genes to the respiratory epithelia of rhesus monkeys. Other instances of the use of adenoviruses in gene therapy can be found in Rosenfeld et al., Science 252:431-434 (1991); Rosenfeld et al., Cell 68:143-155 (1992); Mastrangeli et al., J. Clin. Invest. 91:225-234 (1993); PCT Publication W094/12649; and Wang, et al., Gene Therapy 2:775-783 (1995). In a preferred embodiment, adenovirus vectors are used.

[0519] Adeno-associated virus (AAV) has also been proposed for use in gene therapy (Walsh *et al.*, Proc. Soc. Exp. Biol. Med. 204:289-300 (1993); U.S. Patent No. 5,436,146).

[0520] Another approach to gene therapy involves transferring a gene to cells in tissue culture by such methods as electroporation, lipofection, calcium phosphate mediated transfection, or viral infection. Usually, the method of transfer includes the transfer of a selectable marker to the cells. The cells are then placed under selection to isolate those cells that have taken up and are expressing the transferred gene. Those cells are then delivered to a patient.

[0521] In this embodiment, the nucleic acid is introduced into a cell prior to administration in vivo of the resulting recombinant cell. Such introduction can be carried out by any method known in the art, including but not limited to transfection, electroporation, microinjection, infection with a viral or bacteriophage vector containing the nucleic acid sequences, cell fusion, chromosome-mediated gene transfer, microcell-mediated gene transfer, spheroplast fusion, etc. Numerous techniques are known in the art for the introduction of foreign genes into cells (see, e.g., Loeffler and Behr, Meth.

Enzymol. 217:599-618 (1993); Cohen et al., Meth. Enzymol. 217:618-644 (1993); Clin. Pharma. Ther. 29:69-92m (1985)) and may be used in accordance with the present invention, provided that the necessary developmental and physiological functions of the recipient cells are not disrupted. The technique should provide for the stable transfer of the nucleic acid to the cell, so that the nucleic acid is expressible by the cell and preferably heritable and expressible by its cell progeny.

[0522] The resulting recombinant cells can be delivered to a patient by various methods known in the art. Recombinant blood cells (e.g., hematopoietic stem or progenitor cells) are preferably administered intravenously. The amount of cells envisioned for use depends on the desired effect, patient state, etc., and can be determined by one skilled in the art.

[0523] Cells into which a nucleic acid can be introduced for purposes of gene therapy encompass any desired, available cell type, and include but are not limited to epithelial cells, endothelial cells, keratinocytes, fibroblasts, muscle cells, hepatocytes; blood cells such as T lymphocytes, B lymphocytes, monocytes, macrophages, neutrophils, eosinophils, megakaryocytes, granulocytes; various stem or progenitor cells, in particular hematopoietic stem or progenitor cells, e.g., as obtained from bone marrow, umbilical cord blood, peripheral blood, fetal liver, etc.

[0524] In a preferred embodiment, the cell used for gene therapy is autologous to the patient.

[0525] In an embodiment in which recombinant cells are used in gene therapy, nucleic acid sequences encoding an antibody or fragment thereof are introduced into the cells such that they are expressible by the cells or their progeny, and the recombinant cells are then administered in vivo for therapeutic effect. In a specific embodiment, stem or progenitor cells are used. Any stem and/or progenitor cells which can be isolated and maintained *in vitro* can potentially be used in accordance with this embodiment of the present invention (see e.g. PCT Publication WO 94/08598; Stemple and Anderson, Cell 7 1:973-985 (1992); Rheinwald, Meth. Cell Bio. 21A:229 (1980); and Pittelkow and Scott, Mayo Clinic Proc. 61:771 (1986)).

[0526] In a specific embodiment, the nucleic acid to be introduced for purposes of gene therapy comprises an inducible promoter operably linked to the coding region, such

that expression of the nucleic acid is controllable by controlling the presence or absence of the appropriate inducer of transcription.

Demonstration of Therapeutic or Prophylactic Utility of a Composition

[0527] The compounds of the invention are preferably tested *in vitro*, and then *in vivo* for the desired therapeutic or prophylactic activity, prior to use in humans. For example, *in vitro* assays which can be used to determine whether administration of a specific antibody or composition of the present invention is indicated, include *in vitro* cell culture assays in which a patient tissue sample is grown in culture, and exposed to or otherwise administered an antibody or composition of the present invention, and the effect of such an antibody or composition of the present invention upon the tissue sample is observed. In various specific embodiments, *in vitro* assays can be carried out with representative cells of cell types involved in a patient's disorder, to determine if an antibody or composition of the present invention has a desired effect upon such cell types. Preferably, the antibodies or compositions of the invention are also tested in *in vitro* assays and animal model systems prior to administration to humans.

[0528] Antibodies or compositions of the present invention for use in therapy can be tested for their toxicity in suitable animal model systems, including but not limited to rats, mice, chicken, cows, monkeys, and rabbits. For *in vivo* testing of an antibody or composition's toxicity any animal model system known in the art may be used.

[0529] Efficacy in treating or preventing viral infection may be demonstrated by detecting the ability of an antibody or composition of the invention to inhibit the replication of the virus, to inhibit transmission or prevent the virus from establishing itself in its host, or to prevent, ameliorate or alleviate the symptoms of disease a progression. The treatment is considered therapeutic if there is, for example, a reduction in viral load, amelioration of one or more symptoms, or a decrease in mortality and/or morbidity following administration of an antibody or composition of the invention.

[0530] Antibodies or compositions of the invention can be tested for the ability to induce the expression of cytokines such as IFN-γ, by contacting cells, preferably human cells, with an antibody or composition of the invention or a control antibody or control composition and determining the ability of the antibody or composition of the invention to induce one or more cytokines. Techniques known to those of skill in the art can be used to

measure the level of expression of cytokines. For example, the level of expression of cytokines can be measured by analyzing the level of RNA of cytokines by, for example, RT-PCR and Northern blot analysis, and by analyzing the level of cytokines by, for example, immunoprecipitation followed by western blot analysis and ELISA. In a preferred embodiment, a compound of the invention is tested for its ability to induce the expression of IFN-γ_

Antibodies or compositions of the invention can be tested for their ability to [0531] modulate the biological activity of immune cells by contacting immune cells, preferably human immune cells (e.g., T-cells, B-cells, and Natural Killer cells), with an antibody or composition of the invention or a control compound and determining the ability of the antibody or composition of the invention to modulate (i.e, increase or decrease) the biological activity of immune cells. The ability of an antibody or composition of the invention to modulate the biological activity of immune cells can be assessed by detecting the expression of antigens, detecting the proliferation of immune cells (i.e., B-cell proliferation), detecting the activation of signaling molecules, detecting the effector function of immune cells, or detecting the differentiation of immune cells. Techniques known to those of skill in the art can be used for measuring these activities. For example, cellular proliferation can be assayed by ³H-thymidine incorporation assays and trypan blue cell counts. Antigen expression can be assayed, for example, by immunoassays including, but not limited to, competitive and non-competitive assay systems using techniques such as western blots, immunohistochemistry radioimmunoassays, ELISA (enzyme linked immunosorbent assay), "sandwich" immunoassays, immunoprecipitation assays, precipitin reactions, gel diffusion precipitin reactions, immunodiffusion assays, agglutination assays, complement-fixation assays, immunoradiometric assays, fluorescent immunoassays, protein A immunoassays and FACS analysis. The activation of signaling molecules can be assayed, for example, by kinase assays and electrophoretic shift assays (EMSAs). In a preferred embodiment, the ability of an antibody or composition of the invention to induce B-cell proliferation is measured. In another preferred embodiment, the ability of an antibody or composition of the invention to modulate immunoglobulin expression is measured.

[0532] Antibodies or compositions of the invention can be tested for their ability to reduce tumor formation in *in vitro*, ex vivo and *in vivo* assays. Antibodies or compositions

of the invention can also be tested for their ability to inhibit viral replication or reduce viral load in in vitro and in vivo assays. Antibodies or compositions of the invention can also be tested for their ability to reduce bacterial numbers in in vitro and in vivo assays known to those of skill in the art. Antibodies or compositions of the invention can also be tested for their ability to alleviate of one or more symptoms associated with cancer, an immune disorder (e.g., an inflammatory disease), a neurological disorder or an infectious disease. Antibodies or compositions of the invention can also be tested for their ability to decrease the time course of the infectious disease. Further, antibodies or compositions of the invention can be tested for their ability to increase the survival period of animals suffering from disease or disorder, including cancer, an immune disorder or an infectious disease. Techniques known to those of skill in the art can be used to analyze the function of the antibodies or compositions of the invention in vivo.

Therapeutic/Prophylactic Compositions and Administration

[0533] The invention provides methods of treatment, inhibition and prophylaxis by administration to a subject of an effective amount of antibody (or fragment or variant thereof) or pharmaceutical composition of the invention, preferably an antibody of the invention. In a preferred aspect, an antibody or fragment or variant thereof is substantially purified (i.e., substantially free from substances that limit its effect or produce undesired side-effects). The subject is preferably an animal, including but not limited to, animals such as cows, pigs, horses, chickens, cats, dogs, etc., and is preferably a mammal, and most preferably a human.

[0534] Formulations and methods of administration that can be employed when the compound comprises a nucleic acid or an immunoglobulin are described above; additional appropriate formulations and routes of administration can be selected from among those described herein below.

[0535] Various delivery systems are known and can be used to administer antibody or fragment or variant thereof of the invention, e.g., encapsulation in liposomes, microparticles, microcapsules, recombinant cells capable of expressing the antibody or antibody fragment, receptor-mediated endocytosis (see, e.g., Wu and Wu, J. Biol. Chem. 262:4429-4432 (1987)), construction of a nucleic acid as part of a retroviral or other vector, etc. Methods of introduction include, but are not limited to, intradermal,

intramuscular, intraperitoneal, intravenous, subcutaneous, intranasal, epidural, and oral routes. The compositions may be administered by any convenient route, for example by infusion or bolus injection, by absorption through epithelial or mucocutaneous linings (e.g., oral mucosa, rectal and intestinal mucosa, etc.) and may be administered together with other biologically active agents. Administration can be systemic or local. In addition, it may be desirable to introduce the pharmaceutical compositions of the invention into the central nervous system by any suitable route, including intraventricular and intrathecal injection; intraventricular injection may be facilitated by an intraventricular catheter, for example, attached to a reservoir, such as an Ommaya reservoir. Pulmonary administration can also be employed, e.g., by use of an inhaler or nebulizer, and formulation with an aerosolizing agent.

[0536] In a specific embodiment, it may be desirable to administer the pharmaceutical compositions of the invention locally to the area in need of treatment; this may be achieved by, for example, and not by way of limitation, local infusion during surgery, topical application, e.g., in conjunction with a wound dressing after surgery, by injection, by means of a catheter, by means of a suppository, or by means of an implant, said implant being of a porous, non-porous, or gelatinous material, including membranes, such as sialastic membranes, or fibers. Preferably, when administering a protein, including an antibody, of the invention, care must be taken to use materials to which the protein does not absorb.

[0537] In another embodiment, the composition can be delivered in a vesicle, in particular a liposome (see Langer, Science 249:1527-1533 (1990); Treat *et al.*, in Liposomes in the Therapy of Infectious Disease and Cancer, Lopez-Berestein and Fidler (eds.), Liss, New York, pp. 353-365 (1989); Lopez-Berestein, ibid., pp. 3 17-327; see generally ibid.).

[0538] In yet another embodiment, the composition can be delivered in a controlled release system. In one embodiment, a pump may be used (see Langer, supra; Sefton, CRC Crit. Ref. Biomed. Eng. 14:20 1 (1987); Buchwald *et al.*, Surgery 88:507 (1980); Saudek *et al.*, N. Engl. J. Med. 321:574 (1989)). In another embodiment, polymeric materials can be used (see Medical Applications of Controlled Release, Langer and Wise (eds.), CRC Pres., Boca Raton, Florida (1974); Controlled Drug Bioavailability, Drug Product Design and Performance, Smolen and Ball (eds.), Wiley, New York (1984);

Ranger and Peppas, J., Macromol. Sci. Rev. Macromol. Chem. 23:61 (1983); see also Levy et al., Science 228:190 (1985); During et al., Ann. Neurol. 25:35 1 (1989); Howard et al., J.Neurosurg. 7 1:105 (1989)). In yet another embodiment, a controlled release system can be placed in proximity of the therapeutic target, i.e., the brain, thus requiring only a fraction of the systemic dose (see, e.g., Goodson, in Medical Applications of Controlled Release, supra, vol. 2, pp. 115-138 (1984)).

[0539] Other controlled release systems are discussed in the review by Langer (Science 249:1527-1533 (1990)).

[0540] In a specific embodiment where the composition of the invention is a nucleic acid encoding a protein, the nucleic acid can be administered in vivo to promote expression of its encoded protein, by constructing it as part of an appropriate nucleic acid expression vector and administering it so that it becomes intracellular, e.g., by use of a retroviral vector (see U.S. Patent No. 4,980,286), or by direct injection, or by use of microparticle bombardment (e.g., a gene gun; Biolistic, Dupont), or coating with lipids or cell-surface receptors or transfecting agents, or by administering it in linkage to a homeobox-like peptide which is known to enter the nucleus (see e.g., Joliot et al., Proc. Natl. Acad. Sci. USA 88:1864-1868 (1991)), etc. Alternatively, a nucleic acid can be introduced intracellularly and incorporated within host cell DNA for expression, by homologous recombination.

[0541] The present invention also provides pharmaceutical compositions. Such compositions comprise a therapeutically effective amount of an antibody or a fragment thereof, and a pharmaceutically acceptable carrier. In a specific embodiment, the term "pharmaceutically acceptable" means approved by a regulatory agency of the Federal or a state government or listed in the U.S. Pharmacopeia or other generally recognized pharmacopeia for use in animals, and more particularly in humans. The term "carrier" refers to a diluent, adjuvant, excipient, or vehicle with which the therapeutic is administered. Such pharmaceutical carriers can be sterile liquids, such as water and oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral oil, sesame oil and the like. Water is a preferred carrier when the pharmaceutical composition is administered intravenously. Saline solutions and aqueous dextrose and glycerol solutions can also be employed as liquid carriers, particularly for injectable solutions. Suitable pharmaceutical excipients include starch, glucose, lactose,

sucrose, gelatin, malt, rice, flour, chalk, silica gel, sodium stearate, glycerol monostearate, talc, sodium chloride, dried skim milk, glycerol, propylene, glycol, water, ethanol and the like. The composition, if desired, can also contain minor amounts of wetting or emulsifying agents, or pH buffering agents. These compositions can take the form of solutions, suspensions, emulsion, tablets, pills, capsules, powders, sustained-release formulations and the like. The composition can be formulated as a suppository, with traditional binders and carriers such as triglycerides. Oral formulation can include standard carriers such as pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate, etc. Examples of suitable pharmaceutical carriers are described in "Remington's Pharmaceutical Sciences" by E.W. Martin. Such compositions will contain a therapeutically effective amount of the antibody or fragment thereof, preferably in purified form, together with a suitable amount of carrier so as to provide the form for proper administration to the patient. The formulation should suit the mode of administration.

[0542] In a preferred embodiment, the composition is formulated in accordance with routine procedures as a pharmaceutical composition adapted for intravenous administration to human beings. Typically, compositions for intravenous administration are solutions in sterile isotonic aqueous buffer. Where necessary, the composition may also include a solubilizing agent and a local anesthetic such as lignocamne to ease pain at the site of the injection. Generally, the ingredients are supplied either separately or mixed together in unit dosage form, for example, as a dry lyophilized powder or water free concentrate in a hermetically sealed container such as an ampoule or sachette indicating the quantity of active agent. Where the composition is to be administered by infusion, it can be dispensed with an infusion bottle containing sterile pharmaceutical grade water or saline. Where the composition is administered by injection, an ampoule of sterile water for injection or saline can be provided so that the ingredients may be mixed prior to administration.

[0543] The compositions of the invention can be formulated as neutral or salt forms. Pharmaceutically acceptable salts include those formed with anions such as those derived from hydrochloric, phosphoric, acetic, oxalic, tartaric acids, etc., and those formed with cations such as those derived from sodium, potassium, ammonium, calcium, ferric hydroxides, isopropylamine, triethylamine, 2-ethylamino ethanol, histidine, procaine, etc.

The amount of the composition of the invention which will be effective in the treatment, inhibition and prevention of a disease or disorder associated with aberrant expression and/or activity of a polypeptide of the invention can be determined by standard clinical techniques. In addition, in vitro assays may optionally be employed to help identify optimal dosage ranges. The precise dose to be employed in the formulation will also depend on the route of administration, and the seriousness of the disease or disorder, and should be decided according to the judgment of the practitioner and each patient's circumstances. Effective doses may be extrapolated from dose-response curves derived from in vitro or animal model test systems.

[0545] For antibodies, the dosage administered to a patient is typically 0.1 mg/kg to 100 mg/kg of the patient's body weight. Preferably, the dosage administered to a patient is between 0.1 mg/kg and 20 mg/kg of the patient's body weight, more preferably 1 mg/kg to 10 mg/kg of the patient's body weight. Generally, human antibodies have a longer half-life within the human body than antibodies from other species due to the immune response to the foreign polypeptides. Thus, lower dosages of human antibodies and less frequent administration is often possible. Further, the dosage and frequency of administration of therapeutic or pharmaceutical compositions of the invention may be reduced by enhancing uptake and tissue penetration (e.g., into the brain) of the antibodies by modifications such as, for example, lipidation.

[0546] The antibodies and antibody compositions of the invention may be administered alone or in combination with other adjuvants. Adjuvants that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, alum, alum plus deoxycholate (ImmunoAg), MTP-PE (Biocine Corp.), QS21 (Genentech, Inc.), BCG, and MPL. In a specific embodiment, antibody and antibody compositions of the invention are administered in combination with alum. In another specific embodiment, antibody and antibody compositions of the invention are administered in combination with QS-21. Further adjuvants that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, Monophosphoryl lipid immunomodulator, AdjuVax 100a, QS-21, QS-18, CRL1005, Aluminum salts, MF-59, and Virosomal adjuvant technology. Vaccines that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, vaccines directed toward protection against MMR (measles, mumps,

rubella), polio, varicella, tetanus/diptheria, hepatitis A, hepatitis B, haemophilus influenzae B, whooping cough, pneumonia, influenza, Lyme's Disease, rotavirus, cholera, yellow fever, Japanese encephalitis, poliomyelitis, rabies, typhoid fever, and pertussis, and/or PNEUMOVAX-23™. Combinations may be administered either concomitantly, e.g., as an admixture, separately but simultaneously or concurrently; or sequentially. This includes presentations in which the combined agents are administered together as a therapeutic mixture, and also procedures in which the combined agents are administered separately but simultaneously, e.g., as through separate intravenous lines into the same individual. Administration "in combination" further includes the separate administration of one of the compounds or agents given first, followed by the second.

In another specific embodiment, antibody and antibody compositions of the [0547] invention are used in combination with PNEUMOVAX-23™ to treat, prevent, and/or diagnose infection and/or any disease, disorder, and/or condition associated therewith. In one embodiment, antibody and antibody compositions of the invention are used in combination with PNEUMOVAX-23™ to treat, prevent, and/or diagnose any Gram positive bacterial infection and/or any disease, disorder, and/or condition associated therewith. In another embodiment, antibody and antibody compositions of the invention are used in combination with PNEUMOVAX-23™ to treat, prevent, and/or diagnose infection and/or any disease, disorder, and/or condition associated with one or more members of the genus Enterococcus and/or the genus Streptococcus. In another embodiment, antibody and antibody compositions of the invention are used in any combination with PNEUMOVAX-23™ to treat, prevent, and/or diagnose infection and/or any disease, disorder, and/or condition associated with one or more members of the Group B streptococci. In another embodiment, antibody and antibody compositions of the invention are used in combination with PNEUMOVAX-23™ to treat, prevent, and/or diagnose infection and/or any disease, disorder, and/or condition associated with Streptococcus pneumoniae.

[0548] The antibody and antibody compositions of the invention may be administered alone or in combination with other therapeutic agents, including but not limited to, chemotherapeutic agents, antibiotics, antivirals, steroidal and non-steroidal anti-inflammatories, conventional immunotherapeutic agents and cytokines.

Combinations may be administered either concomitantly, e.g., as an admixture, separately

but simultaneously or concurrently; or sequentially. This includes presentations in which the combined agents are administered together as a therapeutic mixture, and also procedures in which the combined agents are administered separately but simultaneously, e.g., as through separate intravenous lines into the same individual. Administration "in combination" further includes the separate administration of one of the compounds or agents given first, followed by the second.

In one embodiment, the antibody and antibody compositions of the [0549] invention are administered in combination with other members of the TNF family. TNF, TNF-related or TNF-like molecules that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, soluble forms of TNF-alpha, lymphotoxin-alpha (LT-alpha, also known as TNF-beta), LT-beta (found in complex heterotrimer LT-alpha2-beta), OPGL, FasL, CD27L, CD30L, CD40L, 4-1BBL, DcR3, OX40L, TNF-gamma (International Publication No. WO 96/14328), TRAIL, AIM-II (International Publication No. WO 97/34911), APRIL (J. Exp. Med. 188(6):1185-1190 (1998)), endokine-alpha (International Publication No. WO 98/07880), Neutrokine-alpha (International Application Publication No. WO 98/18921), OPG, OX40, and nerve growth factor (NGF), and soluble forms of Fas, CD30, CD27, CD40 and 4-IBB, TR2 (International Publication No. WO 96/34095), DR3 (International Publication No. WO 97/33904), DR4 (International Publication No. WO 98/32856), TR5 (International Publication No. WO 98/30693), TR6 (International Publication No. WO 98/30694), TR7 (International Publication No. WO 98/41629), TRANK, TR9 (International Publication No. WO 98/56892), 312C2 (International Publication No. WO 98/06842), and TR12, and soluble forms CD154, CD70, and CD153.

[0550] In a preferred embodiment, the antibody and antibody compositions of the invention are administered in combination with CD40 ligand (CD40L), a soluble form of CD40L (e.g., AVREND™), biologically active fragments, variants, or derivatives of CD40L, anti-CD40L antibodies (e.g., agonistic or antagonistic antibodies), and/or anti-CD40 antibodies (e.g., agonistic or antagonistic antibodies).

[0551] In an additional embodiment, the antibody and antibody compositions of the invention are administered alone or in combination with an anti-angiogenic agent(s). Anti-angiogenic agents that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, Angiostatin (Entremed,

Rockville, MD), Troponin-1 (Boston Life Sciences, Boston, MA), anti-Invasive Factor, retinoic acid and derivatives thereof, paclitaxel (Taxol), Suramin, Tissue Inhibitor of Metalloproteinase-1, Tissue Inhibitor of Metalloproteinase-2, VEGI, Plasminogen Activator Inhibitor-1, Plasminogen Activator Inhibitor-2, and various forms of the lighter "d group" transition metals.

[0552] Lighter "d group" transition metals include, for example, vanadium, molybdenum, tungsten, titanium, niobium, and tantalum species. Such transition metal species may form transition metal complexes. Suitable complexes of the above-mentioned transition metal species include oxo transition metal complexes.

[0553] Representative examples of vanadium complexes include oxo vanadium complexes such as vanadate and vanadyl complexes. Suitable vanadate complexes include metavanadate and orthovanadate complexes such as, for example, ammonium metavanadate, sodium metavanadate, and sodium orthovanadate. Suitable vanadyl complexes include, for example, vanadyl acetylacetonate and vanadyl sulfate including vanadyl sulfate hydrates such as vanadyl sulfate mono- and trihydrates.

[0554] Representative examples of tungsten and molybdenum complexes also include oxo complexes. Suitable oxo tungsten complexes include tungstate and tungsten oxide complexes. Suitable tungstate complexes include ammonium tungstate, calcium tungstate, sodium tungstate dihydrate, and tungstic acid. Suitable tungsten oxides include tungsten (IV) oxide and tungsten (VI) oxide. Suitable oxo molybdenum complexes include molybdate, molybdenum oxide, and molybdenyl complexes. Suitable molybdate complexes include ammonium molybdate and its hydrates, sodium molybdate and its hydrates, and potassium molybdate and its hydrates. Suitable molybdenum oxides include molybdenum (VI) oxide, molybdenum (VI) oxide, and molybdic acid. Suitable molybdenyl complexes include, for example, molybdenyl acetylacetonate. Other suitable tungsten and molybdenum complexes include hydroxo derivatives derived from, for example, glycerol, tartaric acid, and sugars.

[0555] A wide variety of other anti-angiogenic factors may also be utilized within the context of the present invention. Representative examples include, but are not limited to, platelet factor 4; protamine sulphate; sulphated chitin derivatives (prepared from queen crab shells), (Murata et al., Cancer Res. 51:22-26, 1991); Sulphated Polysaccharide Peptidoglycan Complex (SP-PG) (the function of this compound may be enhanced by the

presence of steroids such as estrogen, and tamoxifen citrate); Staurosporine; modulators of matrix metabolism, including for example, proline analogs, cishydroxyproline, d,L-3,4-dehydroproline, Thiaproline, alpha,alpha-dipyridyl, aminopropionitrile fumarate; 4-propyl-5-(4-pyridinyl)-2(3H)-oxazolone; Methotrexate; Mitoxantrone; Heparin; Interferons; 2 Macroglobulin-serum; ChIMP-3 (Pavloff et al., J. Bio. Chem. 267:17321-17326, 1992); Chymostatin (Tomkinson et al., Biochem J. 286:475-480, 1992); Cyclodextrin Tetradecasulfate; Eponemycin; Camptothecin; Fumagillin (Ingber et al., Nature 348:555-557, 1990); Gold Sodium Thiomalate ("GST"; Matsubara and Ziff, J. Clin. Invest. 79:1440-1446, 1987); anticollagenase-serum; alpha2-antiplasmin (Holmes et al., J. Biol. Chem. 262(4):1659-1664, 1987); Bisantrene (National Cancer Institute); Lobenzarit disodium (N-(2)-carboxyphenyl-4- chloroanthronilic acid disodium or "CCA"; (Takeuchi et al., Agents Actions 36:312-316, 1992); and metalloproteinase inhibitors such as BB94.

[0556] Additional anti-angiogenic factors that may also be utilized within the context of the present invention include Thalidomide, (Celgene, Warren, NJ); Angiostatic steroid; AGM-1470 (H. Brem and J. Folkman J Pediatr. Surg. 28:445-51 (1993)); an integrin alpha v beta 3 antagonist (C. Storgard et al., J Clin. Invest. 103:47-54 (1999)); carboxynaminolmidazole; Carboxyamidotriazole (CAI) (National Cancer Institute, Bethesda, MD); Conbretastatin A-4 (CA4P) (OXiGENE, Boston, MA); Squalamine (Magainin Pharmaceuticals, Plymouth Meeting, PA); TNP-470, (Tap Pharmaceuticals, Deerfield, IL); ZD-0101 AstraZeneca (London, UK); APRA (CT2584); Benefin, Byrostatin-1 (SC339555); CGP-41251 (PKC 412); CM101; Dexrazoxane (ICRF187); DMXAA; Endostatin; Flavopridiol; Genestein; GTE; ImmTher; Iressa (ZD1839); Octreotide (Somatostatin); Panretin; Penacillamine; Photopoint; PI-88; Prinomastat (AG-3340) Purlytin; Suradista (FCE26644); Tamoxifen (Nolvadex); Tazarotene; Tetrathiomolybdate; Xeloda (Capecitabine); and 5-Fluorouracil.

[0557] Anti-angiogenic agents that may be administered in combination with the compounds of the invention may work through a variety of mechanisms including, but not limited to, inhibiting proteolysis of the extracellular matrix, blocking the function of endothelial cell-extracellular matrix adhesion molecules, by antagonizing the function of angiogenesis inducers such as growth factors, and inhibiting integrin receptors expressed on proliferating endothelial cells. Examples of anti-angiogenic inhibitors that interfere

with extracellular matrix proteolysis and which may be administered in combination with the antibody and antibody compositions of the invention include, but are not limited to, AG-3340 (Agouron, La Jolla, CA), BAY-12-9566 (Bayer, West Haven, CT), BMS-275291 (Bristol Myers Squibb, Princeton, NJ), CGS-27032A (Novartis, East Hanover, NJ), Marimastat (British Biotech, Oxford, UK), and Metastat (Aeterna, St-Foy, Quebec). Examples of anti-angiogenic inhibitors that act by blocking the function of endothelial cell-extracellular matrix adhesion molecules and which may be administered in combination with the antibody and antibody compositions of the invention include, but are not limited to, EMD-121974 (Merck KcgaA Darmstadt, Germany) and Vitaxin (Ixsys, La Jolla, CA/Medimmune, Gaithersburg, MD). Examples of anti-angiogenic agents that act by directly antagonizing or inhibiting angiogenesis inducers and which may be administered in combination with the antibody and antibody compositions of the invention include, but are not limited to, Angiozyme (Ribozyme, Boulder, CO), Anti-VEGF antibody (Genentech, S. San Francisco, CA), PTK-787/ZK-225846 (Novartis, Basel, Switzerland), SU-101 (Sugen, S. San Francisco, CA), SU-5416 (Sugen/Pharmacia Upjohn, Bridgewater, NJ), and SU-6668 (Sugen). Other anti-angiogenic agents act to indirectly inhibit angiogenesis. Examples of indirect inhibitors of angiogenesis which may be administered in combination with the antibody and antibody compositions of the invention include, but are not limited to, IM-862 (Cytran, Kirkland, WA), Interferonalpha, IL-12 (Roche, Nutley, NJ), and Pentosan polysulfate (Georgetown University, Washington, DC).

[0558] In particular embodiments, the use of antibody and antibody compositions of the invention in combination with anti-angiogenic agents is contemplated for the treatment, prevention, and/or amelioration of an autoimmune disease, such as for example, an autoimmune disease described herein.

[0559] In a particular embodiment, the use of antibody and antibody compositions of the invention in combination with anti-angiogenic agents is contemplated for the treatment, prevention, and/or amelioration of arthritis. In a more particular embodiment, the use of antibody and antibody compositions of the invention in combination with anti-angiogenic agents is contemplated for the treatment, prevention, and/or amelioration of rheumatoid arthritis.

[0560] In another embodiment, antibody and antibody compositions of the

invention are administered in combination with an anticoagulant. Anticoagulants that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, heparin, warfarin, and aspirin. In a specific embodiment, antibody and antibody compositions of the invention are administered in combination with heparin and/or warfarin. In another specific embodiment, antibody and antibody compositions of the invention are administered in combination with warfarin. In another specific embodiment, antibody and antibody compositions of the invention are administered in combination with warfarin and aspirin. In another specific embodiment, antibody and antibody compositions of the invention are administered in combination with heparin. In another specific embodiment, antibody and antibody compositions of the invention are administered in combination with heparin and aspirin.

[0561] In another embodiment, antibody and antibody compositions of the invention are administered in combination with an agent that suppresses the production of anticardiolipin antibodies. In specific embodiments, the polynucleotides of the invention are administered in combination with an agent that blocks and/or reduces the ability of anticardiolipin antibodies to bind phospholipid-binding plasma protein beta 2-glycoprotein I (b2GPI).

[0562] In certain embodiments, antibody and antibody compositions of the invention are administered in combination with antiretroviral agents, nucleoside reverse transcriptase inhibitors, non-nucleoside reverse transcriptase inhibitors, and/or protease inhibitors. Nucleoside reverse transcriptase inhibitors that may be administered in combination with the antibody and antibody compositions of the invention, include, but are not limited to, RETROVIR™ (zidovudine/AZT), VIDEX™ (didanosine/ddI), HIVID™ (zalcitabine/ddC), ZERIT™ (stavudine/d4T), EPIVIR™ (lamivudine/3TC), and COMBIVIR™ (zidovudine/lamivudine). Non-nucleoside reverse transcriptase inhibitors that may be administered in combination with the antibody and antibody compositions of the invention, include, but are not limited to, VIRAMUNE™ (nevirapine), RESCRIPTOR™ (delayirdine), and SUSTIVA™ (efavirenz). Protease inhibitors that may be administered in combination with the antibody and antibody compositions of the invention, include, but are not limited to, CRIXIVAN™ (indinavir), NORVIR™ (ritonavir), INVIRASE™ (saquinavir), and VIRACEPT™ (nelfinavir). In a specific embodiment, antiretroviral agents, nucleoside reverse transcriptase inhibitors, non-

nucleoside reverse transcriptase inhibitors, and/or protease inhibitors may be used in any combination with antibody and antibody compositions of the invention to treat, prevent, and/or diagnose AIDS and/or to treat, prevent, and/or diagnose HIV infection.

[0563] In other embodiments, antibody and antibody compositions of the invention may be administered in combination with anti-opportunistic infection agents. Antiopportunistic agents that may be administered in combination with the antibody and antibody compositions of the invention, include, but are not limited to, TRIMETHOPRIM-SULFAMETHOXAZOLE™, DAPSONE™, PENTAMIDINE™, ATOVAQUONE™, ISONIAZID™, RIFAMPIN™, PYRAZINAMIDE™, ETHAMBUTOL™, RIFABUTIN™, CLARITHROMYCIN™, AZITHROMYCIN™ GANCICLOVIR™, FOSCARNET™, CIDOFOVIR™, FLUCONAZOLE™. ITRACONAZOLE™, KETOCONAZOLE™, ACYCLOVIR™, FAMCICOLVIR™. PYRIMETHAMINE™, LEUCOVORIN™, NEUPOGEN™ (filgrastim/G-CSF), and LEUKINE™ (sargramostim/GM-CSF). In a specific embodiment, antibody and antibody compositions of the invention are used in any combination with TRIMETHOPRIM-SULFAMETHOXAZOLE™, DAPSONE™, PENTAMIDINE™, and/or ATOVAQUONE™ to prophylactically treat, prevent, and/or diagnose an opportunistic Pneumocystis carinii pneumonia infection. In another specific embodiment, antibody and antibody compositions of the invention are used in any combination with ISONIAZID™, RIFAMPIN™, PYRAZINAMIDE™, and/or ETHAMBUTOL™ to prophylactically treat. prevent, and/or diagnose an opportunistic Mycobacterium avium complex infection. In another specific embodiment, antibody and antibody compositions of the invention are used in any combination with RIFABUTIN™, CLARITHROMYCIN™, and/or AZITHROMYCIN™ to prophylactically treat, prevent, and/or diagnose an opportunistic Mycobacterium tuberculosis infection. In another specific embodiment, antibody and antibody compositions of the invention are used in any combination with GANCICLOVIR™, FOSCARNET™, and/or CIDOFOVIR™ to prophylactically treat, prevent, and/or diagnose an opportunistic cytomegalovirus infection. In another specific embodiment, antibody and antibody compositions of the invention are used in any combination with FLUCONAZOLE™, ITRACONAZOLE™, and/or KETOCONAZOLE™ to prophylactically treat, prevent, and/or diagnose an opportunistic

fungal infection. In another specific embodiment, antibody and antibody compositions of the invention are used in any combination with ACYCLOVIR™ and/or FAMCICOLVIR™ to prophylactically treat, prevent, and/or diagnose an opportunistic herpes simplex virus type I and/or type II infection. In another specific embodiment, antibody and antibody compositions of the invention are used in any combination with PYRIMETHAMINE™ and/or LEUCOVORIN™ to prophylactically treat, prevent, and/or diagnose an opportunistic *Toxoplasma gondii* infection. In another specific embodiment, antibody and antibody compositions of the invention are used in any combination with LEUCOVORIN™ and/or NEUPOGEN™ to prophylactically treat, prevent, and/or diagnose an opportunistic bacterial infection.

[0564] In a further embodiment, the antibody and antibody compositions of the invention are administered in combination with an antiviral agent. Antiviral agents that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, acyclovir, ribavirin, amantadine, and remantidine.

[0565] In a further embodiment, the antibody and antibody compositions of the invention are administered in combination with an antibiotic agent. Antibiotic agents that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, amoxicillin, aminoglycosides, beta-lactam (glycopeptide), beta-lactamases, Clindamycin, chloramphenicol, cephalosporins, ciprofloxacin, ciprofloxacin, erythromycin, fluoroquinolones, macrolides, metronidazole, penicillins, quinolones, rifampin, streptomycin, sulfonamide, tetracyclines, trimethoprim, trimethoprim-sulfamthoxazole, and vancomycin.

[0566] Conventional nonspecific immunosuppressive agents, that may be administered in combination with the antibody and antibody compositions of the invention include, but are not limited to, steroids, cyclosporine, cyclosporine analogs cyclophosphamide, cyclophosphamide IV, methylprednisolone, prednisolone, azathioprine, FK-506, 15-deoxyspergualin, and other immunosuppressive agents that act by suppressing the function of responding T cells.

[0567] In specific embodiments, antibody and antibody compositions of the invention are administered in combination with immunosuppressants.

Immunosuppressants preparations that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, ORTHOCLONETM

(OKT3), SANDIMMUNE™/NEORAL™/SANGDYA™ (cyclosporin), PROGRAF™ (tacrolimus), CELLCEPT™ (mycophenolate), Azathioprine, glucorticosteroids, and RAPAMUNE™ (sirolimus). In a specific embodiment, immunosuppressants may be used to prevent rejection of organ or bone marrow transplantation.

[0568] In a preferred embodiment, the antibody and antibody compositions of the invention are administered in combination with steroid therapy. Steroids that may be administered in combination with the antibody and antibody compositions of the invention, include, but are not limited to, oral corticosteroids, prednisone, and methylprednisolone (e.g., IV methylprednisolone). In a specific embodiment, antibody and antibody compositions of the invention are administered in combination with prednisone. In a further specific embodiment, the antibody and antibody compositions of the invention are administered in combination with prednisone and an immunosuppressive agent. Immunosuppressive agents that may be administered with the antibody and antibody compositions of the invention and prednisone are those described herein, and include, but are not limited to, azathioprine, cylophosphamide, and cyclophosphamide IV. In a another specific embodiment, antibody and antibody compositions of the invention are administered in combination with methylprednisolone. In a further specific embodiment, the antibody and antibody compositions of the invention are administered in combination with methylprednisolone and an immunosuppressive agent. Immunosuppressive agents that may be administered with the antibody and antibody compositions of the invention and methylprednisolone are those described herein, and include, but are not limited to, azathioprine, cylophosphamide, and cyclophosphamide IV.

[0569] In a preferred embodiment, the antibody and antibody compositions of the invention are administered in combination with an antimalarial. Antimalarials that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, hydroxychloroquine, chloroquine, and/or quinacrine.

[0570] In a preferred embodiment, the antibody and antibody compositions of the invention are administered in combination with an NSAID.

[0571] In a nonexclusive embodiment, the antibody and antibody compositions of the invention are administered in combination with one, two, three, four, five, ten, or more of the following drugs: NRD-101 (Hoechst Marion Roussel), diclofenac

(Dimethaid), oxaprozin potassium (Monsanto), mecasermin (Chiron), T-614 (Toyama), pemetrexed disodium (Eli Lilly), atreleuton (Abbott), valdecoxib (Monsanto), eltenac (Byk Gulden), campath, AGM-1470 (Takeda), CDP-571 (Celltech Chiroscience), CM-101 (CarboMed), ML-3000 (Merckle), CB-2431 (KS Biomedix), CBF-BS2 (KS Biomedix), IL-1Ra gene therapy (Valentis), JTE-522 (Japan Tobacco), paclitaxel (Angiotech), DW-166HC (Dong Wha), darbufelone mesylate (Warner-Lambert), soluble TNF receptor 1 (synergen; Amgen), IPR-6001 (Institute for Pharmaceutical Research). trocade (Hoffman-La Roche), EF-5 (Scotia Pharmaceuticals), BIIL-284 (Boehringer Ingelheim), BIIF-1149 (Boehringer Ingelheim), LeukoVax (Inflammatics), MK-663 (Merck), ST-1482 (Sigma-Tau), and butixocort propionate (WarnerLambert). In a preferred embodiment, the antibody and antibody compositions of the [0572] invention are administered in combination with one, two, three, four, five or more of the following drugs: methotrexate, sulfasalazine, sodium aurothiomalate, auranofin, cyclosporine, penicillamine, azathioprine, an antimalarial drug (e.g., as described herein), cyclophosphamide, chlorambucil, gold, ENBREL™ (Etanercept), anti-TNF antibody, LJP 394 (La Jolla Pharmaceutical Company, San Diego, California) and prednisolone. [0573] In a more preferred embodiment, the antibody and antibody compositions of the invention are administered in combination with an antimalarial, methotrexate, anti-TNF antibody, ENBREL™ and/or suflasalazine. In one embodiment, the antibody and antibody compositions of the invention are administered in combination with methotrexate. In another embodiment, the antibody and antibody compositions of the invention are administered in combination with anti-TNF antibody. In another embodiment, the antibody and antibody compositions of the invention are administered in combination with methotrexate and anti-TNF antibody. In another embodiment, the antibody and antibody compositions of the invention are administered in combination with suflasalazine. In another specific embodiment, the antibody and antibody compositions of the invention are administered in combination with methotrexate, anti-TNF antibody, and suflasalazine. In another embodiment, the antibody and antibody compositions of the invention are administered in combination ENBREL™. In another embodiment, the antibody and antibody compositions of the invention are administered in combination with ENBREL™ and methotrexate. In another embodiment, the antibody and antibody compositions of the invention are administered in combination with ENBRELT,

methotrexate and suflasalazine. In another embodiment, the antibody and antibody compositions of the invention are administered in combination with ENBREL™, methotrexate and suflasalazine. In other embodiments, one or more antimalarials is combined with one of the above-recited combinations. In a specific embodiment, the antibody and antibody compositions of the invention are administered in combination with an antimalarial (e.g., hydroxychloroquine), ENBREL™, methotrexate and suflasalazine. In another specific embodiment, the antibody and antibody compositions of the invention are administered in combination with an antimalarial (e.g., hydroxychloroquine), sulfasalazine, anti-TNF antibody, and methotrexate.

In an additional embodiment, antibody and antibody compositions of the invention are administered alone or in combination with one or more intravenous immune globulin preparations. Intravenous immune globulin preparations that may be administered with the antibody and antibody compositions of the invention include, but not limited to, GAMMAR™, IVEEGAM™, SANDOGLOBULIN™, GAMMAGARD S/D™, and GAMIMUNE™. In a specific embodiment, antibody and antibody compositions of the invention are administered in combination with intravenous immune globulin preparations in transplantation therapy (e.g., bone marrow transplant).

[0575] CD40 ligand (CD40L), a soluble form of CD40L (e.g., AVREND™), biologically active fragments, variants, or derivatives of CD40L, anti-CD40L antibodies (e.g., agonistic or antagonistic antibodies), and/or anti-CD40 antibodies (e.g., agonistic or antagonistic antibodies).

[0576] In an additional embodiment, the antibody and antibody compositions of the invention are administered alone or in combination with an anti-inflammatory agent. Anti-inflammatory agents that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, glucocorticoids and the nonsteroidal anti-inflammatories, aminoarylcarboxylic acid derivatives, arylacetic acid derivatives, arylbutyric acid derivatives, arylcarboxylic acids, arylpropionic acid derivatives, pyrazoles, pyrazolones, salicylic acid derivatives, thiazinecarboxamides, eacetamidocaproic acid, S-adenosylmethionine, 3-amino-4-hydroxybutyric acid, amixetrine, bendazac, benzydamine, bucolome, difenpiramide, ditazol, emorfazone, guaiazulene, nabumetone, nimesulide, orgotein, oxaceprol, paranyline, perisoxal, pifoxime, proquazone, proxazole, and tenidap.

[0577]In another embodiment, compostions of the invention are administered in combination with a chemotherapeutic agent. Chemotherapeutic agents that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, antibiotic derivatives (e.g., doxorubicin, bleomycin, daunorubicin, and dactinomycin); antiestrogens (e.g., tamoxifen); antimetabolites (e.g., fluorouracil, 5-FU, methotrexate, floxuridine, interferon alpha-2b, glutamic acid, plicamycin, mercaptopurine, and 6-thioguanine); cytotoxic agents (e.g., carmustine, BCNU, lomustine, CCNU, cytosine arabinoside, cyclophosphamide, estramustine, hydroxyurea, procarbazine, mitomycin, busulfan, cis-platin, and vincristine sulfate); hormones (e.g., medroxyprogesterone, estramustine phosphate sodium, ethinyl estradiol, estradiol, megestrol acetate, methyltestosterone, diethylstilbestrol diphosphate, chlorotrianisene. and testolactone); nitrogen mustard derivatives (e.g., mephalen, chorambucil, mechlorethamine (nitrogen mustard) and thiotepa); steroids and combinations (e.g., bethamethasone sodium phosphate); and others (e.g., dicarbazine, asparaginase, mitotane, vincristine sulfate, vinblastine sulfate, and etoposide).

[0578] In a specific embodiment, antibody and antibody compositions of the invention are administered in combination with CHOP (cyclophosphamide, doxorubicin, vincristine, and prednisone) or any combination of the components of CHOP. In another embodiment, antibody and antibody compositions of the invention are administered in combination with Rituximab. In a further embodiment, antibody and antibody compositions of the invention are administered with Rituximab and CHOP, or Rituxmab and any combination of the components of CHOP.

In an additional embodiment, the antibody and antibody compositions of the invention are administered in combination with cytokines. Cytokines that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, GM-CSF, G-CSF, IL2, IL3, IL4, IL5, IL6, IL7, IL10, IL12, IL13, IL15, anti-CD40, CD40L, IFN-alpha, IFN-beta, IFN-gamma, TNF-alpha, and TNF-beta. In preferred embodiments, antibody and antibody compositions of the invention are administered with BLyS (e.g., amino acids 134-285 of SEQ IF D NO:3228). In another embodiment, antibody and antibody compositions of the invention may be administered with any interleukin, including, but not limited to, IL-1alpha, IL-1beta, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, IL-14, IL-15, IL-16, IL-17, IL-18,

IL-19, IL-20, IL-21, and IL-22. In preferred embodiments, the antibody and antibody compositions of the invention are administered in combination with IL4 and IL10.

[0580] In one embodiment, the antibody and antibody compositions of the invention are administered in combination with one or more chemokines. In specific embodiments, the antibody and antibody compositions of the invention are administered in combination with an $\alpha(CxC)$ chemokine selected from the group consisting of gammainterferon inducible protein-10 (yIP-10), interleukin-8 (IL-8), platelet factor-4 (PF4), neutrophil activating protein (NAP-2), GRO-α, GRO-β, GRO-γ, neutrophil-activating peptide (ENA-78), granulocyte chemoattractant protein-2 (GCP-2), and stromal cellderived factor-1 (SDF-1, or pre-B cell stimulatory factor (PBSF)); and/or a β(CC) chemokine selected from the group consisting of: RANTES (regulated on activation, normal T expressed and secreted), macrophage inflammatory protein-1 alpha (MIP-1α), macrophage inflammatory protein-1 beta (MIP-1β), monocyte chemotactic protein-1 (MCP-1), monocyte chemotactic protein-2 (MCP-2), monocyte chemotactic protein-3 (MCP-3), monocyte chemotactic protein-4 (MCP-4) macrophage inflammatory protein-1 gamma (MIP-1γ), macrophage inflammatory protein-3 alpha (MIP-3α), macrophage inflammatory protein-3 beta (MIP-3β), macrophage inflammatory protein-4 (MIP-4/DC-CK-1/PARC), eotaxin, Exodus, and I-309; and/or the γ(C) chemokine, lymphotactin.

[0581] In another embodiment, the antibody and antibody compositions of the invention are administered with chemokine beta-8, chemokine beta-1, and/or macrophage inflammatory protein-4. In a preferred embodiment, the antibody and antibody compositions of the invention are administered with chemokine beta-8.

In an additional embodiment, the antibody and antibody compositions of the invention are administered in combination with an IL-4 antagonist. IL-4 antagonists that may be administered with the antibody and antibody compositions of the invention include, but are not limited to: soluble IL-4 receptor polypeptides, multimeric forms of soluble IL-4 receptor polypeptides; anti-IL-4 receptor antibodies that bind the IL-4 receptor without transducing the biological signal elicited by IL-4, anti-IL4 antibodies that block binding of IL-4 to one or more IL-4 receptors, and muteins of IL-4 that bind IL-4 receptors but do not transduce the biological signal elicited by IL-4. Preferably, the antibodies employed according to this method are monoclonal antibodies (including antibody fragments, such as, for example, those described herein).

The invention also encompasses combining the polynucleotides and/or [0583] polypeptides of the invention (and/or agonists or antagonists thereof) with other proposed or conventional hematopoietic therapies. Thus, for example, the polynucleotides and/or polypeptides of the invention (and/or agonists or antagonists thereof) can be combined with compounds that singly exhibit erythropoietic stimulatory effects, such as erythropoietin, testosterone, progenitor cell stimulators, insulin-like growth factor, prostaglandins, serotonin, cyclic AMP, prolactin, and triiodothyzonine. Also encompassed are combinations of the antibody and antibody compositions of the invention with compounds generally used to treat aplastic anemia, such as, for example, methenolene, stanozolol, and nandrolone; to treat iron-deficiency anemia, such as, for example, iron preparations; to treat malignant anemia, such as, for example, vitamin B₁₂ and/or folic acid; and to treat hemolytic anemia, such as, for example, adrenocortical steroids, e.g., corticoids. See e.g., Resegotti et al., Panminerva Medica, 23:243-248 (1981); Kurtz, FEBS Letters, 14a:105-108 (1982); McGonigle et al., Kidney Int., 25:437-444 (1984); and Pavlovic-Kantera, Expt. Hematol., 8(supp. 8) 283-291 (1980), the contents of each of which are hereby incorporated by reference in their entireties. Compounds that enhance the effects of or synergize with erythropoietin are also useful as adjuvants herein, and include but are not limited to, adrenergic agonists, thyroid hormones, androgens, hepatic erythropoietic factors, erythrotropins, and erythrogenins, See for e.g., Dunn, "Current Concepts in Erythropoiesis", John Wiley and Sons (Chichester, England, 1983); Kalmani, Kidney Int., 22:383-391 (1982); Shahidi, New Eng. J. Med., 289:72-80 (1973); Urabe et al., J. Exp. Med., 149:1314-1325 (1979); Billat et al., Expt. Hematol., 10:133-140 (1982); Naughton et al., Acta Haemat, 69:171-179 (1983); Cognote et al. in abstract 364, Proceedings 7th Intl. Cong. of Endocrinology (Quebec City, Quebec, July 1-7, 1984); and Rothman et al., 1982, J. Surg. Oncol., 20:105-108 (1982). Methods for stimulating hematopoiesis comprise administering a hematopoietically effective amount (i.e., an amount which effects the formation of blood cells) of a pharmaceutical composition containing polynucleotides and/or poylpeptides of the invention (and/or agonists or antagonists thereof) to a patient. The polynucleotides and/or polypeptides of the invention and/or agonists or antagonists thereof is administered to the patient by any suitable technique, including but not limited to, parenteral, sublingual, topical, intrapulmonary and intranasal, and those techniques further discussed

herein. The pharmaceutical composition optionally contains one or more members of the group consisting of erythropoietin, testosterone, progenitor cell stimulators, insulin-like growth factor, prostaglandins, serotonin, cyclic AMP, prolactin, triiodothyzonine, methenolene, stanozolol, and nandrolone, iron preparations, vitamin B₁₂, folic acid and/or adrenocortical steroids.

[0585] In an additional embodiment, the antibody and antibody compositions of the invention are administered in combination with hematopoietic growth factors. Hematopoietic growth factors that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, LEUKINE™ (SARGRAMOSTIM™) and NEUPOGEN™ (FILGRASTIM™).

[0586] In an additional embodiment, the antibody and antibody compositions of the invention are administered in combination with fibroblast growth factors. Fibroblast growth factors that may be administered with the antibody and antibody compositions of the invention include, but are not limited to, FGF-1, FGF-2, FGF-3, FGF-4, FGF-5, FGF-6, FGF-7, FGF-8, FGF-9, FGF-10, FGF-11, FGF-12, FGF-13, FGF-14, and FGF-15.

[0587] Additionally, the antibody and antibody compositions of the invention may be administered alone or in combination with other therapeutic regimens, including but not limited to, radiation therapy. Such combinatorial therapy may be administered sequentially and/or concomitantly.

Kits .

[0588] The invention also provides a pharmaceutical pack or kit comprising one or more containers filled with one or more of the ingredients of the pharmaceutical compositions of the invention. Optionally associated with such container(s) can be a notice in the form prescribed by a governmental agency regulating the manufacture, use or sale of pharmaceuticals or biological products, which notice reflects approval by the agency of manufacture, use or sale for human administration.

[0589] The present invention provides kits that can be used in the above methods. In one embodiment, a kit comprises an antibody of the invention, preferably a purified antibody, in one or more containers. In an alternative embodiment, a kit comprises an antibody fragment that immunospecifically binds to BLyS. In a specific embodiment, the kits of the present invention contain a substantially isolated BLyS polypeptide as a control.

Preferably, the kits of the present invention further comprise a control antibody which does not react with BLyS. In another specific embodiment, the kits of the present invention contain a means for detecting the binding of an antibody to BLyS (e.g., the antibody may be conjugated to a detectable substrate such as a fluorescent compound, an enzymatic substrate, a radioactive compound or a luminescent compound, or a second antibody which recognizes the first antibody may be conjugated to a detectable substrate). In specific embodiments, the kit may include a recombinantly produced or chemically synthesized BLyS. The BLyS provided in the kit may also be attached to a solid support. In a more specific embodiment the detecting means of the above-described kit includes a solid support to which BLyS is attached. Such a kit may also include a non-attached reporter-labeled anti-human antibody. In this embodiment, binding of the antibody to BLyS can be detected by binding of the said reporter-labeled antibody.

[0590] In an additional embodiment, the invention includes a diagnostic kit for use in screening serum containing antigens of the polypeptide of the invention. The diagnostic kit includes a substantially isolated antibody specifically immunoreactive with BLyS, and means for detecting the binding of BLyS to the antibody. In one embodiment, the antibody is attached to a solid support. In a specific embodiment, the antibody may be a monoclonal antibody. The detecting means of the kit may include a second, labeled monoclonal antibody. Alternatively, or in addition, the detecting means may include a labeled, competing antigen.

[0591] In one diagnostic configuration, test serum is reacted with a solid phase reagent having a surface-bound BLyS obtained by the methods of the present invention. After BLyS binds to a specific antibody, the unbound serum components are removed by washing, reporter-labeled anti-human antibody is added, unbound anti-human antibody is removed by washing, and a reagent is reacted with reporter-labeled anti-human antibody to bind reporter to the reagent in proportion to the amount of bound anti-BLyS antibody on the solid support. Typically, the reporter is an enzyme which is detected by incubating the solid phase in the presence of a suitable fluorometric, luminescent or colorimetric substrate.

[0592] The solid surface reagent in the above assay is prepared by known techniques for attaching protein material to solid support material, such as polymeric beads, dip sticks, 96-well plate or filter material. These attachment methods generally

include non-specific adsorption of the protein to the support or covalent attachment of the protein, typically through a free amine group, to a chemically reactive group on the solid support, such as an activated carboxyl, hydroxyl, or aldehyde group. Alternatively, streptavidin coated plates can be used in conjunction with biotinylated antigen(s).

[0593] Thus, the invention provides an assay system or kit for carrying out this diagnostic method. The kit generally includes a support with surface-bound recombinant BLyS, and a reporter-labeled anti-human antibody for detecting surface-bound anti-BLyS antibody.

[0594] In specific embodiments, the present invention encompasses a single chain Fv (scFv) having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0595] In specific embodiments, the present invention encompasses a single chain Fv (scFv) having an amino acid sequence of one of SEQ ID NOS: 1 to 46, 321 to 329, 1563 to 1595, and 1881 to 1908.

[0596] In specific embodiments, the present invention encompasses a single chain Fv (scFv) having an amino acid sequence of one of SEQ ID NOS: 1563 to 1880.

[0597] In specific embodiments, the present invention encompasses a single chain Fy (scFy) having an amino acid sequence of one of SEQ ID NOS: 1881 to 2128.

[0598] In specific embodiments, the present invention encompasses a single chain Fv (scFv) having an amino acid sequence of one of SEQ ID NOS: 1 to 1562.

[0599] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS.

[0600] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 46, 321 to 329, 1563 to 1595, and 1881 to 1908.

[0601] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1881 to 2128, and in which said antibody or fragment thereof immunospecifically binds to the membrane-bound form of BLyS.

[0602] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1563 to 1880, and in which said antibody or fragment thereof immunospecifically binds to the soluble form of BLyS.

[0603] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS.

[0604] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 46, 321 to 329, 1563 to 1595, and 1881 to 1908.

[0605] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1881 to 2128, and in which said antibody or fragment thereof immunospecifically binds to the membrane-bound form of BLyS.

[0606] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1563 to 1880, and in which said antibody or fragment thereof immunospecifically binds to the soluble form of BLyS.

[0607] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and which also comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0608] In specific embodiments, the present invention encompasses an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and which also comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.and in which said VL and said VH domains are derived from the same scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0609] In specific embodiments, the present invention encompasses an antibody or

fragment thereof comprising an amino acid sequence of one of SEQ ID NOS: 2129 to 3227 wherein said antibody or fragment thereof immunospecifically binds BLyS.

[0610] In specific embodiments, the antibody or fragment thereof of the invention is a whole immunoglobulin molecule.

[0611] In specific embodiments, the antibody or fragment thereof of the invention is a Fab fragment.

[0612] In specific embodiments, the antibody or fragment thereof of the invention is a Fv fragment.

[0613] In specific embodiments, the present invention encompasses a chimeric protein comprising the antibody or fragment thereof of the invention covalently linked to a heterologous polypeptide.

[0614] In specific embodiments, the present invention encompasses a composition comprising two or more types of antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VH domain from a different scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0615] In specific embodiments, the present invention encompasses a composition comprising two or more types of antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VL domain from a different scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0616] In specific embodiments, the present invention encompasses a composition comprising two or more types of antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VL domain from a different scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128 and wherein each type of antibody or fragment thereof further comprises a VH domain from a different scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0617] In specific embodiments, the present invention encompasses a composition comprising two or more types of antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VH CDR3 having an amino acid sequence of one of SEQ ID

NOS: 3129 to 3227.

[0618] In specific embodiments, the present invention encompasses a panel of two or more types of antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VH domain from a different scFv having an amino acid sequence of one of SEQ ID NO: 1 to 2128.

[0619] In specific embodiments, the present invention encompasses a panel of two or more types of antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VL domain from a different scFv having an amino acid sequence of one of SEQ ID NO: 1 to 2128.

[0620] In specific embodiments, the present invention encompasses a panel of two or more types of antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VL domain from a different scFv having an amino acid sequence of one of SEQ ID NO: 1 to 2128 and wherein each type of antibody or fragment further comprises a VH domain from a different scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0621] In specific embodiments, the present invention encompasses a panel of two or more antibodies or fragments or variants thereof, each of which type immunospecifically binds to BLyS, and each of which type of antibody or fragment thereof comprises a VHCDR3 from a different scFv having an amino acid sequence of one of SEQ ID NOS: 2129 to 3227.

[0622] In specific embodiments, the antibodies or fragments thereof of the antibody panel of the invention, are each in a well of a 96 well plate.

[0623] In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS.

[0624] In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment

thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 46, 321 to 329, 1563 to 1595, and 1881 to 1908, wherein said antibody or fragment thereof immunospecifically binds BLyS.

[0625] In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1881 to 1908, wherein the antibody of fragment thereof immunospecifically binds the membrane-bound form of BLyS.

In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1563 to 1569, wherein said antibody of fragment thereof immunospecifically binds the soluble form of BLyS. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host

cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 46, 321 to 329, 1563 to 1595, and 1881 to 1908, wherein said antibody or fragment thereof immunospecifically binds BLyS. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1881 to 2128, wherein the antibody of fragment thereof immunospecifically binds the membrane-bound form of BLyS. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

[0630] In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1563 to 1880, wherein said antibody of fragment thereof immunospecifically binds the soluble form of BLyS. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

[0631] In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and which also comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

[0632] In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VL domain from an scFv having an amino acid sequence of one of

SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and which also comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128 and in which said VL domain and said VH domain are derived from the same scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

In specific embodiments, the present invention encompasses an isolated nucleic acid molecule comprising a nucleotide sequence encoding an antibody or fragment thereof comprising a VHCDR3 from an scFv having an amino acid sequence of one of SEQ ID NOS: 2129 to 3227, wherein said antibody or fragment thereof immunospecifically binds BLyS. The present invention also encompasses vectors comprising the isolated nucleic acid molecule described above, including vectors comprising a nucleotide sequence which regulates the expression of the antibody or fragment thereof encoded by the above-described nucleic acid molecule. Additionally the present invention also encompasses host cells, including mammalian host cells, comprising the above-described nucleic acid molecule which is operably linked to a heterologous promoter, as well as host cells, including mammalian host cells, comprising the above-described vectors. Additionally, the present invention also provides a method for producing an antibody or fragment thereof comprising culturing the above-described host cells under conditions in which the nucleic acid molecule is expressed.

[0634] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment thereof comprising an amino acid sequence of a VH domain encoded by a nucleotide sequence that hybridizes under stringent conditions to a nucleotide sequence encoding a

VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0635] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment thereof comprising an amino acid sequence of a VL domain encoded by a nucleotide sequence that hybridizes under stringent conditions to a nucleotide sequence encoding a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0636] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment thereof comprising an amino acid sequence of a VH domain encoded by a nucleotide sequence that hybridizes under highly stringent conditions to a nucleotide sequence encoding a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0637] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment thereof comprising an amino acid sequence of a VL domain encoded by a nucleotide sequence that hybridizes under highly stringent conditions to a nucleotide sequence encoding a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0638] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment thereof comprising an amino acid sequence of a CDR encoded by a nucleotide sequence that hybridizes under stringent conditions to a nucleotide sequence encoding a CDR from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0639] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment thereof comprising an amino acid sequence of a CDR encoded by a nucleotide sequence that hybridizes under highly stringent conditions to a nucleotide sequence encoding a CDR from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0640] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment

thereof comprising an amino acid sequence of a VH CDR3 encoded by a nucleotide sequence that hybridizes under stringent conditions to a nucleotide sequence encoding a VH CDR3 having an amino acid sequence of one of SEQ ID NOS: 2129 to 3227.

[0641] In specific embodiments, the present invention provides an antibody or fragment thereof that immunospecifically binds to BLyS, said antibody or fragment thereof comprising an amino acid sequence of a VH CDR3 encoded by a nucleotide sequence that hybridizes under highly stringent conditions to a nucleotide sequence encoding a VH CDR3 having an amino acid sequence of one of SEQ ID NOS: 2129 to 3227.

[0642] In specific embodiments, the present invention provides a method for detecting of aberrant expression of BLyS, comprising:

[0643] assaying the level of BLyS expression in cells or a tissue sample of an individual using one or more antibodies or fragments or variants thereof that immunospecifically bind BLyS; and

[0644] comparing the level of BLyS assayed in the cells or a tissue sample with a standard level of BLyS or a level of BLyS in cells or a tissue sample from an individual without aberrant BLyS expression, wherein an increase or decrease in the assayed level of BLyS or level in cells or a tissue sample from an individual without aberrant BLyS expression compared to the standard level of BLyS is indicative of aberrant expression.

[0645] In specific embodiments, the present invention provides a method for diagnosing a disease or disorder associated with aberrant BLyS expression or activity, comprising:

[0646] administering to a subject an effective amount of a labeled antibody or fragment thereof that immunospecifically binds to BLyS;

[0647] waiting for a time interval following the administering for permitting the labeled antibody or fragment thereof to preferentially concentrate at sites in the subject where BLyS is expressed;

[0648] determining background level; and

[0649] detecting the labeled antibody or fragment thereof in the subject, such that detection of labeled antibody or fragment thereof above the background level indicates that the subject has a particular disease or disorder associated with aberrant expression of BLyS.

[0650] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0651] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above comprises a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128.

[0652] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above comprises a VH CDR3 having an amino acid sequence of one of SEQ ID NOS: 2129 to 3227.

[0653] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above is conjugated to a diagnostic agent.

[0654] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above is conjugated to a diagnostic agent wherein the diagnostic agent is horseradish peroxidase, alkaline phosphatase, beta-galactosidase, or acetylcholinesterase.

[0655] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above is conjugated to a diagnostic agent wherein the diagnostic agent is fluorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine fluorescein, dansyl chloride or phycoerythrin.

[0656] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above is conjugated to a diagnostic agent wherein the diagnostic agent is ¹²⁵I, ¹³¹I, ¹¹¹In, ⁹⁰Y or ⁹⁹Tc.

[0657] In specific embodiments, the antibody or fragment thereof utilized in the two methods described immediately above is conjugated to a diagnostic agent wherein the diagnostic agent is luciferase, luciferin or aequorin.

[0658] A pharmaceutical composition comprising at least one antibody or fragment thereof of comprising a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and a pharmaceutically acceptable carrier.

[0659] A pharmaceutical composition comprising at least one antibody or fragment thereof of comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof

immunospecifically binds BLyS and a pharmaceutically acceptable carrier.

[0660] A pharmaceutical composition comprising at least one antibody or fragment thereof of comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and which also comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128 and a pharmaceutically acceptable carrier.

[0661] A pharmaceutical composition comprising at least one antibody or fragment thereof of comprising an amino acid sequence of one of SEQ ID NOS: 2129 to 3227 wherein said antibody or fragment thereof immunospecifically binds BLyS and a pharmaceutically acceptable carrier.

[0662] A method of treating, preventing or ameliorating a disease or disorder associated with aberrant BLyS expression or activity, comprising administering to an animal in need thereof the pharmaceutical composition comprising at least one antibody or fragment thereof of comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and which also comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128 and a pharmaceutically acceptable carrier in an amount effective to treat, prevent or ameliorate the disease or disorder. This method may be used to treat an infectious disorder, cancer, and/or an autoimmune disease such as lupus or glomerular nephritis.

[0663] A method of treating, preventing or ameliorating a disease or disorder associated with aberrant BLyS expression or activity, comprising administering to an animal in need thereof the pharmaceutical composition comprising at least one antibody or fragment thereof of comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and a pharmaceutically acceptable carrier in an amount effective to treat, prevent or ameliorate the disease or disorder. This method may be used to treat an infectious disorder, cancer, and/or an autoimmune disease such as lupus or glomerular nephritis.

[0664] A method of treating, preventing or ameliorating a disease or disorder associated with aberrant BLyS expression or activity, comprising administering to an

animal in need thereof the pharmaceutical composition comprising at least one antibody or fragment thereof of comprising a VL domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128, wherein said antibody or fragment thereof immunospecifically binds BLyS and which also comprises a VH domain from an scFv having an amino acid sequence of one of SEQ ID NOS: 1 to 2128 and a pharmaceutically acceptable carrier in an amount effective to treat, prevent or ameliorate the disease or disorder. This method may be used to treat an infectious disorder, cancer, and/or an autoimmune disease such as lupus or glomerular nephritis.

[0665] A method of treating, preventing or ameliorating a disease or disorder associated with aberrant BLyS expression or activity, comprising administering to an animal in need thereof the pharmaceutical composition of comprising at least one antibody or fragment thereof of comprising an amino acid sequence of one of SEQ ID NOS: 2129 to 3227 wherein said antibody or fragment thereof immunospecifically binds BLyS and a pharmaceutically acceptable carrier in an amount effective to treat, prevent or ameliorate the disease or disorder. This method may be used to treat an infectious disorder, cancer, and/or an autoimmune disease such as lupus or glomerular nephritis.

[0666] This method may be used to treat an infectious disorder, cancer, and/or an autoimmune disease such as lupus or glomerular nephritis.

EXAMPLES

Abbreviations

0.2 M Tris-HCl, 0.5 mM EDTA, 0.5 M sucrose (TES)

1-ethyl-3-[3-dimethylaminopropyl]carbo diimide hydrochloride (EDC)

2TY supplemented with 100µg/ml ampicillin and 2% glucose (2TYAG)

2TY supplemented with 100µg/ml ampicillin and 50µg/ml kanamycin (2TYAK)

3,3',5,5'-Tetramethyl Benzidine (TMB)

50% inhibitory concentration (IC₅₀)

6xPBS containing 18% Marvel blocking solution (6xMPBS)

Absorbance (A)

Bovine serum albumin (BSA)

Enzyme linked immunosorbent assay (ELISA)

Foetal calf serum (FCS)

Heavy chain variable (V_H)

Hepes buffered saline (HBS)

Horseradish peroxidase (HRP)

Immobilised Metal Affinity Chromatography (IMAC)

Isopropyl B-D-thiogalactopyranoside (IPTG)

Light chain variable(V_L)

Multiplicity of infection (MOI)

N-[2-hydroxyethyl]piperazine-N'-[2-ethanesulfonic acid] (Hepes)

Nanomolar (nM)

N-Hydroxysuccinimide (NHS)

PBS containing 3% Marvel (MPBS)

Phosphate Buffered Saline (PBS)

Phosphate Buffered Saline + 0.1% (v/v) Tween 20 (PBST)

Picomolar (pM)

Single chain fragment variable (scFv)

Tumour Necrosis Factor-alpha (TNF-a)

Tumour Necrosis Factor-beta (TNF-β)

TNF-related apoptosis inducing ligand (TRAIL)

Definitions:

In the following section "immobilized BLyS" refers to a soluble form of BLyS or biotinylated BLyS coated on a plastic assay plate (e.g., a 96 well plate), but does not refer to histidine tagged BLyS coated on a plastic assay plate.; "biotinylated BLyS" is a soluble form of BLyS except when used to coat an ELISA plate, in which case it would be "immobilized BLyS." Membrane bound forms of BLyS include, but are not limited to, U937 and P388 plasma membranes.

Example 1: Antibodies Immunospecifically Binding to Soluble And Membrane-Bound BLyS

displaying scFvs that immunospecifically bind to the soluble and membrane-bound forms of BLyS. Phage displaying scFvs that bound to immobilized BLyS were identified after panning on immobilized BLyS and assessment by ELISA for binding to immobilized BLyS. The BLyS that was immobilized on plates for these assays was purified from supernatants of Sf9 cells infected with a baculovirus expression construct as described in Moore et al., Science 285:260-263 which is hereby incorporated by reference in its entirety. Each of the identified scFvs were then sequenced. Certain sequences were isolated multiple times, thus a panel (panel 1) containing one member of each unique sequences was generated and further characterized for their ability to immunospecifically bind to the soluble and membrane-bound forms of BLyS.

[0669] The derived amino acid sequences of these scFvs are shown in Table 1 above. The individual V_H and V_L segments of the scFvs were aligned to the known human germline sequences in V-BASE (Tomlinson et al, www.mrc-cpe.cam.ac.uk) and the closest germline identified.

Example 2: Specificity of scFvs for BLyS and Membrane-Bound BLyS

[0670] The specificity of each of the scFvs for both BLyS and membrane-bound BLyS was determined by phage ELISA. BLyS was immobilised onto plastic as a purified soluble form of the protein or as a membrane-bound form present on plasma membrane preparations from the human macrophage-like cell line, U937.

Maintenance of U937 Cells

[0671] U937 cells are a human monocyte-like, histiocytic lymphoma cell line known to express BLyS on their plasma membranes. They were maintained in RPMI-1640 supplemented with 4mM L-glutamine, 10% FCS, 10 U penicillin, 100 g/ml streptomycin (all reagents from Sigma). The cells were thawed from frozen stock and are either used for plasma membrane preparation, or split 1:5, after 2 days in culture when the cell density reaches 1 x 10⁶/ml.

Preparation of U937 Plasma Membranes

To prepare plasma membranes, 1x109 U937 cells were harvested from their [0672] culture medium by centrifugation at 1000 rpm at 4°C for 5 minutes in a benchtop centrifuge. The cells were resuspended in 40 ml 12 mM Tris, pH 7.5, 250 mM sucrose and placed on ice. The cells are then lysed using a hand-held electric homogenizer (Labortechnik IKA Ultra-Turrax) for four, one minute, bursts. To check that cell lysis had occurred, 10 µl cell lysate was added to 10 µl Trypan blue and the cell lysate was examined under a microscope. After confirming lysis, the homogenate was centrifuged at 270 x g, for 10 minutes at 4°C to pellet the nuclear fraction and the supernatant was retained. The supernatant was centrifuged at 8000 x g, 10 mins, 4°C, to pellet the mitochondrial and lysosomal fractions and the supernatant was retained. The supernatant was then centrifuged at 100000 x g, 60 mins, 4°C to pellet the plasma membrane enriched fraction. The supernatant was discarded and the plasma membrane pellet was resuspended in 1 ml PBS and stored at -70°C. The protein concentration of the plasma membrane fraction was determined using a protein quantification kit (Biorad). Typical yields were between 5 and 10 mg of plasma membranes.

Phage ELISA

[0673] To determine the specificity of each of the unique scFvs, a phage ELISA was performed for each scFv against human BLyS, U937 plasma membranes, TNFα (R&D Systems, Minneapolis, MN), BSA and uncoated well. Individual *E. coli* colonies containing a phagemid representing one of the unique scFvs from panel 1 were inoculated into 96-well plates containing 100 μl 2TYAG medium per well. Plates were incubated at 37°C for 4 hours, shaking. M13KO7 helper phage was added to each well to a MOI of 10 and the plates were incubated for a further 1 hour at 37°C. The plates were centrifuged in a benchtop centrifuge at 2000 rpm for 10 minutes. The supernatant was removed and cell pellets were resuspended in 100 μl 2TYAK and incubated at 30°C overnight, shaking. The next day, plates were centrifuged at 2000 rpm for 10 min and the 100 μl phage-containing supernatant from each well carefully transferred into a fresh 96-well plate. Twenty μl of 6xMPBS was added to each well, and incubated at room temperature for 1 hour to pre-block the phage prior to ELISA.

[001]Flexible 96-well plates (Falcon) were coated overnight at 4°C with human BLyS (1 µg/ml) in PBS, U937 plasma membranes (10 µg/ml) in PBS, TNFa (1 µg/ml) in PBS, BSA (1 µg/ml) in PBS, or PBS. After coating, the solutions were removed from the wells, and the plates were blocked for 1 hour at room temperature in MPBS. The plates were washed 3 times with PBS and then 50 µl of pre-blocked phage was added to each well. The plates were incubated at room temperature for 1 hour and then washed with 3 changes of PBST followed by 3 changes of PBS. To each well, 50 µl of an anti-gene VIII-HRP conjugate (Pharmacia) at a 1 to 5000 dilution in MPBS was added and the plates incubated at room temperature for 1 hour. Each plate was washed three times with PBST followed by three times with PBS. Then 50 µl of an HRP-labelled anti-mouse polymer (DAKO EnVision) diluted 1/50 in 3% MPBS was added and incubated for 1 hour at room temperature. Each plate was then washed three times with PBST followed by three times with PBS. Fifty µl of TMB substrate was then added to each well, and incubated at room temperature for 30 minutes or until colour development. The reaction was stopped by the addition of 25 µl of 0.5 M H₂SO₄. The signal generated was measured by reading the absorbance at 450nm (A₄₅₀) using a microtiter plate reader (Bio-Rad 3550).

[001] The results for 3 clones (I006E07, I008D05 and I016F04) are shown in Figure 1. All 3 scFvs recognize immobilized BLyS and U937 plasma membranes but do not recognize TNFa, BSA or an uncoated well (PBS only). These results indicate that these scFvs specifically recognize immobilized BLyS and membrane-bound BLyS.

Example 3: Inhibition in an *In Vitro* Receptor Binding Assay by Phage ScFvs [0676] All of the unique phage scFvs in panel 1 were assessed for their ability to inhibit soluble BLyS binding to its cognate receptor on IM9 cells.

Biotinylation of BLyS

[0677] One hundred µg of either human or mouse BLyS was dialysed overnight at 4°C against 50 mM sodium bicarbonate (sodium hydrogen carbonate) pH8.5 using a slide-a-lyzer cassette (Pierce). The next day, NHS-biotin (Pierce) was dissolved in DMSO to 13.3 mg/ml. This was then added to the BLyS at a molar ratio of 20:1 biotin:BLyS, mixed

and incubated on ice for 2 hours. The biotinylated BLyS was then dialysed back into sterile PBS (Sigma) using a slide-a-lyzer cassette overnight at 4°C. The biological activity of the biotinylated BLyS was confirmed using the receptor binding inhibition assay (see below).

Maintenance of IM9 cells

[0678] IM9 cells are a human B lymphocyte cell line. They were maintained in RPMI-1640 supplemented with 4 mM L-glutamine, 10% FCS, 10 U penicillin, 100 g/ml streptomycin (all reagents from Sigma). The cells are thawed from frozen stock and can be used in assays after 5 days in culture when they reach a density of 4 - 8 x 10⁵ /ml.

Receptor binding inhibition assay

Individual E. coli colonies containing a phagemid representing one of the unique scFvs from panel 1 were inoculated into 96-well plates containing 100 µl 2TYAG medium per well. Plates were incubated at 37° C for 4 hours, shaking. M13KO7 helper phage was added to each well to a MOI of 10 and the plates were incubated for a further 1 hour at 37°C. The plates were centrifuged in a benchtop centrifuge at 2000 rpm for 10 minutes. The supernatant was removed and cell pellets were resuspended in 100 µl 2TYAK and incubated at 30°C overnight, shaking. The next day, plates were centrifuged at 2000 rpm for 10 min and the 100 µl phage-containing supernatant from each well carefully transferred into a fresh 96-well plate. Phage were diluted 1 in 2 in MPBS prior to use.

[001] Flat-bottomed 96-well plates (Costar) were coated with 100 µl per well of a 1:10 dilution of poly-L-lysine (Sigma) in PBS for 1 hour at room temperature. The plates were then washed twice with water, allowed to air-dry and placed at 4° C overnight. One hundred µl of IM9 cells (at 10⁶/ml in RPMI-1640 culture medium) were then added to each well. Plates were then centrifuged at 3200 rpm for 5 mins to pellet the cells. The media was carefully aspirated and 200 µl of MPBS added to each well. The plates were then allowed to block for 1 hour at room temperature.

[0681] To a separate 96-well plate 10 µl of biotinylated BLyS (at 162.5 ng/ml) in MPBS was added to each well to give a final concentration of 25 ng/ml. Fifty-five µl of

each appropriate phage supernatant was added to each well and the final volume in each well was 65 µl. Plates were then incubated at room temperature for 30 minutes.

[0682] The IM9 coated plates were washed twice in PBS, tapped dry and immediately 50 µl of the phage/biotinylated-BLyS mix was added and incubated at room temperature for 1 hour. Plates were washed three times in PBST and three times in PBS, tapped dry and 50 µl of streptavidin-Delfia (Wallac) was added to each well at 1:1000 dilution in the Manufacturer's assay buffer. The plates were then incubated at room temperature for 1 hour and washed six times in Delfia wash solution (Wallac). After tapping the plates dry, 100 µl per well of Delfia enhancement solution (Wallac) was added. The plates were gently tapped to encourage micelle formation, incubated at room temperature for 10 minutes, and fluorescence read on a Wallac 1420 workstation at 6520 nM.

[0683] Results for 3 phage scFvs (I001C09, I018D07 and I016H07) that inhibited the binding of biotinylated BLyS are shown in Figure 2. Maximal binding of biotinylated BLyS to its receptor (bio-BLyS only), the background signal in the absence of biotinylated BLyS (no bio-BLyS), and results with an irrelevant (i.e., does not recognize BLyS) phage antibody are also shown. All 3 phage scFvs inhibited biotinylated BLyS binding to its receptor on IM9 cells, identifying these scFvs as scFvs that bind the soluble form of BLyS. These scFvs also bind to U937 membranes, thus they also bind the membrane bound form of BLyS.

[0684] Forty-eight of the scFvs from panel 1 that demonstrated the greatest inhibition as phage particles in this assay were chosen for further study. These 48 scFvs are listed in Table 3.

Table 3. scFvs that Inhibit the Binding of Biotinylated-BLyS to its Receptor

Antibody	Antibody	Antibody	Antibody	Antibody
I008C02	I029D07	I008C03	I008C12	I028A06
I022E02	I061E07	I007H08	I061H01	I031C03
I018C02	I006D07	I008A11	I006D08	I031F02

I008B01	I017D10	I061D02	I026E03	I031F09
I016F04	I007B03	I008A09	I027A07	I031G11
I016E05	I018C10	I007F11	I016H07	1050A07
1018H08	I001C09	I037E07	I021B05	I050A12
I018H09	I018D07	I037E12	I031G10	I050B11
	I029F11	I016F02	I031G08	I051C04
	I022D01		I031C07	I003F12
	<u> </u>		I012A06	

Example 4: Specificity of Anti-BLyS Antibodies

[0685] The specificity of the 48 scFvs listed in Table 3 for human and murine BLyS was determined using phage ELISA.

Phage ELISA

To determine the specificity of the 48 scFvs, a phage ELISA was [001] performed against human and mouse BLyS, and a panel of related and unrelated human antigens: Fas ligand, TRAIL, TNFα, TNFβ, and PBS. The: Fas ligand, TRAIL, TNFα, and TNFB antigens were obtained from R&D Systems, Minneapolis, MN. Individual E. coli colonies containing phagemid were inoculated into 5 ml 2YTAG and incubated at 37°C for 4 hours, shaking. M13KO7 helper phage (Pharmacia) was added to each tube to a MOI of 10 and incubated for 30 minutes at 37°C for 1 hour, the first 30 minutes static and the final 30 minutes with gentle shaking. Cells were pelleted by centrifugation at 3,500 rpm for 10 minutes and the supernatant discarded. Cell pellets were resuspended in 5 ml 2TYAK and incubated at 30°C overnight with shaking. The next day, the cells were pelleted by centrifugation at 3,500 rpm for 10 minutes. The phage-containing supernatant (5 ml) was carefully transferred to a fresh tube, 1 ml of 6MPBS was added, and the tube was incubated at room temperature for 1 hour to pre-block the phage prior to ELISA. All antigens were coated at 1 µg/ml. ELISAs were performed essentially [0687]

labelled anti-mouse polymer was omitted. Binding to mouse BLyS was detected with TMB as in Section Example 2.

[0688] All 48 scFvs are specific for immobilized human BLyS and 43 out of the 48 scFvs cross-react with immobilized mouse BLyS but not with any other unrelated or related antigen tested. I008C03, I007F11, I037E07, I037E12, and I016H07 did not bind murine BLyS. Results for two scFvs, I022D01 and I031F02, are shown in Figure 3. Both these scFvs specifically recognize human and mouse BLyS but not any other unrelated or related antigen tested.

Example 5: Specificity for the Membrane-Bound Form of BLyS

[0689] The specificity of 48 scFvs for membrane-bound BLyS was determined by the phage ELISA described in Example 2. BLyS was immobilised onto plastic as a membrane-bound form present on plasma membranes preparations from the human macrophage-like cell line, U937. This cell line is known to express the membrane-bound form of human BLyS.

[0690] To demonstrate that this binding is specific for membrane-bound BLyS, a competition ELISA was developed to determine if the ELISA signal for an individual antibody on U937's could be competed out by pre-incubation with either BLyS or TNF α . An anti-BLyS antibody that also recognizes membrane-bound BLyS would be expected to demonstrate a signal reduction with free BLyS but not free TNF α .

Competition ELISA

Individual E. coli colonies containing phagemid for each of the 48 scFvs listed in Table 3 were inoculated into 5 ml 2YTAG and incubated at 37°C for 4 hours, shaking. M13KO7 helper phage (Pharmacia) was added to each tube to a MOI of 10 and incubated for 30 minutes at 37° C for 1 hour, the first 30 minutes static and the final 30 minutes with gentle shaking. Cells were pelleted by centrifugation at 3,500 rpm for 10 minutes and the supernatant discarded. Cell pellets were resuspended in 5 ml 2TYAK and incubated at 30°C overnight with shaking. The next day, the cells were pelleted by

centrifugation at 3,500 rpm for 10 minutes. The phage-containing supernatants (5 ml) were carefully transferred to a fresh tube.

[0692] For each of the 48 scFvs listed in Table 3, two aliquots of 20 μl 6xMPBS were pipetted into separate wells of a 96-well plate (Greiner). The first aliquot was supplemented with BLyS to a final concentration of 0.5 μg/ml. The second aliquot was supplemented with TNF-α to a final concentration of 0.5 μg/ml. Each experiment was performed in triplicate. One hundred μl of each phage supernatant was then added to each aliquot and mixed by pipetting up and down. The phage were incubated (± competing antigen) at room temperature for 1 hour.

[0693] Flexible 96-well plates (Falcon) were coated overnight at 4°C with 50 µl of 10 μg/ml U937 plasma membranes. After coating, the plates were washed 3 times with PBS and blocked for 1 hour at room temperature with 200 µl MPBS. The plates were washed 3 times with PBS and 50 µl of phage (± competing antigen) was added to each appropriate well. The plates were incubated at room temperature for 1 hour and then washed with 3 changes of PBST followed by 3 changes of PBS. To each well, 50 µl of a mouse anti-gene VIII-HRP conjugate (Pharmacia) at a 1:5000 dilution in MPBS was added and the plates incubated at room temperature for 1 hour. Each plate was washed three times with PBST followed by three times with PBS. Then 50 µl of an HRP-labelled anti-mouse polymer (DAKO EnVision) diluted 1:50 in 3% MPBS was added and incubated for 1 hour at room temperature. Each plate was then washed three times with PBST followed by three times with PBS. Fifty µl of TMB substrate was then added to each well, and incubated at room temperature for 30 to 60 minutes or until color development. The reaction was stopped by the addition of 25 µl of 0.5 M H₂SO₄. The signal generated was measured by reading the absorbance at 450nm (A_{450}) using a microtiter plate reader (Bio-Rad 3550).

[0694] All 48 scFvs bind to U937 plasma membrane preparations. This signal could be competed out by pre-incubation of the phage antibody with BLyS but not by pre-incubation with TNF- α . This indicates that the 48 scFvs specifically recognize membrane-bound BLyS as well as soluble BLyS. Typical results are exemplified by scFvs I031F09, I050A12 and I051C04 and are shown in Figure 4. All 3 scFvs demonstrate binding to U937 plasma membranes. This binding was specifically competed

out with BLyS but did not compete with TNF- α , demonstrating specific recognition of membrane-bound BLyS.

Example 6: scFv Off-rate Determinations

[0695] All off-rate determinations were performed on BIAcore 2000 machines, using the BIAcore 2000 Control Software and evaluated using the BIAevaluation 3.0 software.

Preparation of a Low Density BLyS Surface

[0696] A 500RU surface was prepared for kinetic studies with purified scFvs. A low density BLyS surface (500 RU BLyS coupled) was prepared in flow cell 2 by amine coupling to a CM5 chip. A new CM5 chip was inserted into the BIAcore and a sensorgram initiated with HBS buffer at a flow rate of 5 µl/min. The NHS and EDC coupling solutions (BIAcore) were mixed according to manufacturer's instructions and 30 µl injected over the CM5 surface. Fifty µl of BLyS at 1 µg/ml in 10 mM sodium acetate buffer, pH4, was then injected followed by 30 µl of ethanolamine-HCl solution (BIAcore). The flow rate was then adjusted to 20 µl/min and 10 µl of 4M guanidine hydrochloride in HBS injected over the surface. This strips the surface of non-covalently bound BLyS.

Measurement of scFv off-rate kinetics on the low density surfaces

[0697] The chip containing the low density BLyS surface was inserted in to the BIAcore. A dilution series of purified scFvs was prepared in HBS, typically 50 μ g/ml doubling dilutions down to 1.5 μ g/ml. The dilution series was then injected sequentially over the low density BLyS surface (and blank control) using the following program:

MAIN

FLOWCELL 1,2,3,4

APROG	genab	rldl	ab1
APROG	genab .	r1d2	ab2
APROG	genab	r1d3	ab3
APROG	genab	r1d4	ab4

APROG genab

rld5 ab5

APROG

genab

rld6 ab6

APPEND CONTINUE

END

DEFINE APROG genab

PARAM %Abpos %AbId

FLOW

20

KINJECT

%Abpos 200 80

INJECT

r1c6 10! guanidine hydrochloride regeneration step

EXTRACLEAN

END

[0698] Bound scFvs were removed by injecting $10\mu l$ 4M GuHCl in HBS over the surface between scFv samples.

[0699] The binding curves for individual scFvs were analyzed using the BIAevaluation software to determine antibody off-rates. Kinetic analysis for a typical scFv antibody, 1003C02, is shown in Figure 5. 1003C02 has a $K_{off} = 6 \times 10^{-3} \text{ s}^{-1}$.

Example 7: Inhibition in an In Vitro Receptor Binding Assay by scFv Antibodies

[0700] The 48 scFvs listed in Table 3 were purified and assessed for their ability to inhibit BLyS binding to its receptor on IM9 cells.

Purification of scFv

[0701] To determine the inhibitory potency of anti-BLyS scFv, scFv's were first prepared by IMAC. 2TYAG (5 ml) was inoculated with a single colony and grown overnight at 30°C, shaking. This overnight culture was then used to inoculate 500 ml of 2TY containing 100 μ g/ml ampicillin and 0.1% Glucose, and grown at 30°C, shaking, until an A₆₀₀ of 1.0 was attained. IPTG was added to 1 mM and the culture was grown for a further 3.5 hours at 30°C.

[0702] Cells were harvested by centrifugation at 5,000rpm, and resuspended in 10 ml of TES. A further 15 ml of a 1:5 dilution (in water) of TES was added, and the cell suspension incubated on a turning wheel at 4°C for 30 minutes. This causes osmotic shock and yields a periplasmic extract containing the scFv. Residual cells and debris were pelleted by centrifugation at 9,000 rpm for 20 minutes at 4°C. The supernatant was transferred to a new tube, and 50 μ l of 1 M MgCl₂ added. Two ml of a Ni-NTA agarose (Qiagen), pre-washed with buffer (50 mM sodium phosphate, pH 8, 300 mM NaCl) together with a protease inhibitor tablet (Boehringer Mannheim) were then added to the periplasmic extract. The preparation was incubated, rotating, overnight at 4°C. The Ni-NTA was pelleted by centrifugation at 2,000 rpm for 5 minutes, and the supernatant was aspirated. The agarose beads were washed 3 times with 50 ml wash buffer, centrifuging to collect the agarose in between each wash. Ten ml of wash buffer was added after the final wash, and the slurry was loaded on to a polyprep column (BioRad). Two ml elution buffer (50 mM NaPi (sodium phosphate), pH 8, 300 mM NaCl, 250 mM imidazole) was added to the drained agarose, and the eluate was collected. IMAC purified scFv was buffer exchanged in to PBS by use of a Nap 5 column (Pharmacia) according to the manufacturer's instructions. The A280 was read and the protein concentration determined using a molar extinction coefficient of 1 mg/ml protein = A_{200} 1.4. Purified scFv was stored in 500 μ l aliquots at -70°C.

Receptor Binding Inhibition Assay

[001] Flat-bottomed 96-well plates (Costar) were coated with 100 µl per well of a 1:10 dilution of poly-L-lysine (Sigma) in PBS for 1 hour at room temperature. The plates were then washed twice with water, allowed to air-dry and placed at 4 °C overnight. One hundred µl of IM9 cells (at 10⁶/ml in RPMI-1640) were then added to each well. Plates were then centrifuged at 3200 rpm for 5 mins to pellet the cells. The media was carefully aspirated and 200 µl of MPBS added to each well. The plates were then left to block for 1 hour at room temperature.

[0704] To a separate 96-well plate, titrate test scFvs in MPBS, in triplicate, over a concentration range from 10 µg/ml down to 0.001 µg/ml were added. The final volume of test scFv in each well was 55 µl. Competition with unlabelled BLyS was also included in every assay as a control. Unlabelled BLyS, in MPBS, was typically titrated in triplicate, over a concentration range from 1 µg/ml down to 0.001 µg/ml. 10 µl of biotinylated-BLyS (at 162.5 ng/ml) in MPBS was added to each well to give a final concentration of 25 ng/ml. Plates were then incubated at room temperature for 30 minutes.

[0705] The IM9 coated plates was washed twice in PBS, tapped dry and immediately 50µl of the scFv/biotinylated-BLyS mix was added and incubated at room temperature for 1 hour. Plates were washed three times in PBST and three times in PBS, tapped dry and 50 µl per well added of streptavidin-Delfia (Wallac) at 1:1000 dilution in the Manufacturer's assay buffer. The plates were then incubated at room temperature for 1 hour and washed six times in Delfia wash solution (Wallac). After tapping the plates dry, 100µl per well of Delfia enhancement solution (Wallac) was added. The plates were gently tapped to encourage micelle formation, incubated at room temperature for 10 minutes, and fluorescence read on a Wallac 1420 workstation at 6520 nM.

[0706] Typical titration curves for two scFv antibodies, I007F11 and I050A07, are shown in Figure 6. Unlabelled BLyS competed for binding to its receptor with an IC $_{50}$ value of 0.8 nM. The IC $_{50}$ values for I007F11 and I050A07 are 7.9 nM and 17.1 nM, respectively. The assay was performed in triplicate and standard error bars are shown. The 9 scFvs that demonstrated the greatest inhibition as scFv are listed in Table 4. This data also confirms that these9 scFvs recognize the soluble form of BLyS.

Table 4: 9 ScFvs that demonstrated greatest potency in BLyS Receptor Binding Inhibition Assay

ScFv Antibody
I017D10
I022D01
I008A11
I006D08

I031F02	
I050A12	
I050B11	
1051C04	
I003F12S	

Example 8: Antibodies recognizing a soluble form of BLyS

[0707] A library of phage was screened in an assay to identify those phage displaying scFvs that immunospecifically bind to the soluble but not the membrane-bound forms of BLyS.

[0708] A phage library was screened for the ability to bind to biotinylated BLyS. The phage were exposed to biotinylated BLyS, allowed an interval of time to bind the biotinylated BLyS. Phage binding bio-BLyS were then isolated by capture on streptavidin coated magnetic beads.

[0709] The phage identified in the screen above (capture of Bio-BLyS from solution) were then screened by ELISA for their ability to bind immobilized BLyS. The scFv expressed by phage that bound immobilized BLyS were then cloned and sequenced. Again, several sequences were identified multiple times, thus a panel (panel 2) consisting of on example of each phage expressing a unique scFv was then characterized further.

[0710] The derived amino acid sequences of these scFvs are shown in Table 1 above. The individual V_H and V_L segments of the scFvs were aligned to the known human germline sequences in V-BASE (Tomlinson et al, www.mrc-cpe.cam.ac.uk) and the closest germline identified.

Example 9: Specificity For Soluble BLyS

[0711] The scFvs were isolated from a library of phage based on their ability to bind a soluble form of BLyS. Briefly, phage were preincubated with biotinylated BLyS in solution. Phage that bound to this biotinylated BLyS were then isolated using streptavidin coated magnetic beads.

[0712] The specificity of each of the unique scFvs for BLyS and for the membrane-bound form of BLyS, was determined by phage ELISA. BLyS was immobilised onto plastic as a purified soluble form of the protein or as a membrane-bound

form present on plasma membrane preparations from the human macrophage-like cell line, U937. Maintenance of U937 cells and plasma membrane preparations were performed as detailed in Example 2.

Phage ELISA

[0713] To determine the specificity of each of the scFvs, a phage ELISA was performed for each antibody against human BLyS, U937 plasma membranes, TNFα, BSA and an uncoated well. Antigen coating conditions were as described in Example 2, apart from human BLyS. BLyS was first biotinylated (as described in Example 3) and coated at 1 μg/ml onto streptavidin coated plates (Reacti-Bind, Pierce) for 30 mins at room temperature. The plates were then washed, blocked and the phage ELISA performed as detailed in Example 2.

[001] The results for 3 clones (I074B12, I075F12 and I075A02) that bind the soluble but not the membrane-bound form of BLyS are shown in Figure 7. As a control, a phage antibody that recognizes TNFα, is also shown in Figure 7. There is a small non-specific background signal on the U937 plasma membranes that is evident with both the anti-BLyS scFvs as well as the anti-TNFα control. All 3 anti-BLyS scFvs recognize BLyS but not U937 plasma membranes, TNFα, BSA or an uncoated well (PBS only). This indicates that the scFvs do not bind the membrane-bound form of BLyS. Further, The fact that these scFvs were isolated on the basis of their ability to bind soluble biotinylated BLyS indicates that they bind the soluble form of BLyS. Further confirmation of these scFvs' specificity for BLyS is provided in Example 10.

Example 10: Inhibition in an *in vitro* receptor binding assay by phage scFvs

[0715] All of the unique phage scFvs from panel 2 were assessed for their ability to inhibit BLyS binding to its cognate receptor on IM9 cells. The biotinylation of BLyS, maintenance of IM9 cells and receptor binding inhibition assay were performed as described in Example 3.

[0716] Results for two phage scFvs, I0025B09 and I026C04 are shown in Figure 8. Maximal binding of biotinylated BLyS to its receptor (bio-BLyS only), the background signal in the absence of biotinylated BLyS (no bio-BLyS), and results with an irrelevant (i.e. does not recognize BLyS) phage antibody are also shown. Both phage scFvs inhibited

biotinylated BLyS binding to its receptor on IM9 cells. 33 of the unique scFvs from panel 2 were identified for further study. These 33 scFvs demonstrated the greatest inhibition as phage particles in this assay and are listed in Table 5.

Table 5: Identification of 33 phage scFvs to free BLyS that demonstrate the most significant inhibition of biotinylated-BLyS binding to its receptor

Antibody	Antibody	Antibody	Antibody
I026C04	I074B12	I073F04	I065D04
I003C06	I075A02	I078D08	I068C08
I025B09	I068B08	I078D02	I068F03
I027B12	I068B04	I075G01	I069B07
I025B06	I068C06	I071B03	·
I030A10	I075F12	I072B09	91
I002A01R	I065D08	1078Н08	
I002A01K	I065F08	I064C04	
I026C04R	I067B10	I064C07	
I026C04K	I067F05		

Example 11: Specificity of anti-BLyS scFvs

[0717] The specificity of the 33 scFvs (listed in Table 5) for immobilized human and murine BLyS was determined using phage ELISA.

Phage ELISA

[001] To determine the specificity of the 33 scFvs, a phage ELISA was performed as described in Example 4 against human and mouse BLyS, and a panel of related human antigens: TRAIL, LIGHT, TNFa, TNFb, and an uncoated well (PBS only).

[0719] Typical results for two scFvs, I067F05 and I078D02 are shown in Figure 9. A control antibody that specifically recognizes TNFa is also shown. Both anti-BLyS scFvs specifically recognize immobilized human and mouse BLyS but not any other antigen tested.

[0720] All 33 scFvs are specific for human BLyS. 14/33 cross-react with mouse BLyS but not with any other unrelated or related antigen tested.

Example 12: scFv Off-Rate Determinations

[0721] Off-rate determinations, preparation of a low density BLyS surface and kinetic measurements were as detailed in Example 6.

[0722] The binding curves for individual scFvs were analysed using the BIAevaluation software to determine antibody off-rates. Kinetic analysis for a typical scFv antibody, I002A01, is shown in Figure 10. I002A01 has a $K_{off} = 9 \times 10^{-4} \text{ s}^{-1}$.

Example 13: Inhibition in an in vitro receptor binding assay by scFv antibodies

[0723] The 33 scFvs identified in Table 5 were prepared as purified scFvs and assessed for their ability to inhibit BLyS binding to its receptor on IM9 cells. The scFvs were purified and analysed in the receptor binding inhibition assay as described in Example 6.1.8.

[0724] Typical titration curves for two scFvs, I0068C06 and I074B12, are shown in Figure 11. Unlabelled BLyS competed for binding to its receptor with an inhibitory constant 50 (IC $_{50}$) value of 0.66 nM. The IC $_{50}$ values for I0068C06 and I074B12 are 61 nM and 13 nM , respectively. The assay was performed in triplicate and standard error bars are shown. The 7 scFvs that demonstrated the greatest inhibition as scFv are listed in Table 6.

Table 6: Identification of 7 scFvs to free BLyS that demonstrate the most significant inhibition of biotinylated-BLyS binding to its receptor as purified scFv's.

Antibody
I002A01-R
I002A01-K
I026C04-R
I026C04-K
I068C06
I075F12
I067B10

Example 14: ScFvs Recognizing Membrane-bound BLyS

[0725] A library of phage was screened in an assay to identify those phage displaying scFvs that immunospecifically bind to the membrane-bound but not the soluble form of BLyS.

[0726] As a starting point, a library of phage expressing scFv antibodies were panned on immobilized HIS-tagged BLyS. Phage isolated by panning were then screened for the ability to bind to HIS-tagged BLyS. HIS-tagged BLyS was obtained by expressing amino acids 71-285 of SEQ ID NO:3228 using the pQE9 vector (Qiagen Inc., Valencia, CA) in *E. coli* and purifying the expressed protein. This phage clones identified by this screen were then sequenced. After sequencing, A panel (panel 3) of phage each expressing a unique scFv that bound HIS-tagged BLyS was generated and further characterized.

[0727] The derived amino acid sequences of the unique scFvs from panel 3 are shown in Table 1 above. The individual V_H and V_L segments of the scFvs were aligned to the known human germline sequences in V-BASE (Tomlinson et al, www.mrc-cpe.cam.ac.uk) and the closest germline identified.

Example 15: Recognition of Membrane-bound BLyS

[0728] The specificity of each of the unique scFvs for both the membrane-bound form of BLyS as well as for the soluble form of BLyS, was determined by phage ELISA.

[0729] BLyS was immobilised onto plastic either directly as a purified soluble form of the protein or biotinylated and coated on a streptavidin plate as in Example 9. Binding to HIS-tagged BLyS was used as a primary screen for scFv's that would bind the membrane-bound form of BLyS (see below). The membrane-bound form of BLyS was presented as plasma membranes preparations from the human macrophage-like cell line, U937 or the murine cell line P388.

[0730] Mouse monoclonal antibodies have been raised against His-tagged BLyS according to standard procedures. Characterization of these mouse monoclonal antibodies revealed that they specifically recognized both His-tagged BLyS and the membrane-bound form of BLyS on U937 cells, but not soluble BLyS. Therefore, specific recognition of His-tagged BLyS was used as supporting evidence for the recognition of the membrane-bound form of BLyS by phage and scFv antibodies.

Phage ELISA

[001] To determine the specificity of each of the scFvs, a phage ELISA was performed for each antibody against His-tagged human BLyS, U937 plasma membranes, TNF α , BSA and an uncoated well. Antigen coating conditions were as described in 2. apart from human BLyS. BLyS was first biotinylated (as described in Example 3) and coated at 1 μ g/ml onto streptavidin coated plates (Reacti-Bind, Pierce) for 30 mins at room temperature. The plates were then washed, blocked and the phage ELISA performed as detailed in Example 2.

[0732] The results for 3 clones, I079C01, I081C10 and I082A02, and a control phage antibody that recognizes TNFα, are shown in Figure 12. All 3 scFvs recognize U937 plasma membranes (U937) and His-tagged BLyS (HIS-BLyS) but not, biotinylated BLyS (bio-BLyS) or an uncoated well (PBS). This indicates that the scFvs recognize the membrane-bound form of BLyS.

Example 16: Specificity for Membrane-bound BLyS

[0733] The specificity of the scFvs for only the membrane-bound form of BLyS, and not for the soluble form, was confirmed using a competition ELISA. This assay assesses the ability of test phage scFvs to bind to the membrane-bound form of BLyS on U937 plasma membranes in the presence of different forms of competing BLyS. Competing BLyS was either the His-tagged form of BLyS or soluble BLyS. ScFvs specific for the membrane-bound BLyS would be expected to be competed out by pre-incubation with His-tagged BLyS but not by pre-incubation with soluble BLyS.

[0734] Maintenance of U937 cells and plasma membrane preparations were performed as detailed in Example 2.

Competition ELISA

[0735] U937 plasma membranes (50µl per well) were coated at 10µg/ml in PBS onto Falcon 96-well plates overnight at 4°C.

[0736] Individual E. coli colonies containing a phagemid representing one of the unique scFvs from the panel 3 were inoculated into 50 ml tubes (Falcon) containing 5 ml 2TYAG medium. Tubes were incubated at 37°C for 4 hours, shaking. M13KO7 helper phage was added to each tube to an MOI of 10 and the tubes were incubated for a further 1

hour at 37°C. The tubes were centrifuged in a benchtop centrifuge at 3500 rpm for 10 minutes. The supernatant was removed and cell pellets were resuspended in 5 ml 2TYAK and incubated at 30°C overnight, shaking. The next day, tubes were centrifuged at 3500 rpm for 10 min and the phage-containing supernatant carefully transferred into a fresh tube.

[0737] For each test phage antibody, 3 aliquots of 20µl 18% marvel/6xPBS were transferred into separate wells of a 96-well plate. The first aliquot was supplemented with His-tagged BLyS to a final concentration of 60 µg/ml. The second aliquot was supplemented with soluble BLyS to a final concentration of 60 µg/ml. The third aliquot was not supplemented with any competing antigen. One hundred µl of phage supernatant was then added to each aliquot and left to block at room temperature for 1 hour.

[0738] The antigen-coated plates were washed once with PBS before the addition of 200 μl/well 3% marvel/PBS. These plates were left to block at 37°C for 1 hour and were then washed once with PBS. Duplicate samples of 50 μl pre-blocked phage (above) were added to the antigen-coated plates and left at room temperature for 1 hour. Plates were washed 3x with PBS/0.1%Tween 20, then 3x with PBS. Fifty μl/well mouse anti-M13 HRP (Pharmacia) at 1/5000 in 3% Marvel/PBS was added and left for 1 hour at room temperature. Plates were washed 3 times with PBS/0.1%Tween 20, then 3 times with PBS. Fifty μl/well HRP-labelled anti-mouse Envision polymer (DAKO) at 1/50 in 3% marvel/PBS was added and left for 1 hour at RT. Plates were washed 3 times with PBS/0.1% Tween 20, then 3 times with PBS. Next, 50μl/well of TMB (Sigma) was added and plates left to develop for 30 to 60 minutes. When sufficient color has developed, 25μl/well 0.5M H₂SO₄ was added to stop the reaction. The plates were read at 450nm on a microtiter plate reader (Bio-Rad 3550).

[001] The results for 3 clones, I079B04, I079F08 and I080B01, and a control phage antibody that recognizes TNFα, are shown in Figure 13. All 3 scFvs recognize U937 plasma membranes (U937). This binding is competed out to background levels (i.e. comparable to the signal observed with the anti-TNFα phage antibody) in the presence of His-tagged BLyS (HIS-BLyS) but not biotinylated BLyS (bio-BLyS). This confirms that the scFvs specifically recognize the membrane-bound form but not the soluble form of BLyS.

Example 17: High Throughput BIAcore Screen to identify high affinity scFvs

[0740] This is a 96-well screen where the test samples (scFvs) are derived from 1 ml periplasmic extracts of individual antibody expressing clones. Potentially higher affinity scFvs are then identified principally as those giving a large number of total RU's bound to a HIS-BLyS surface in BIAcore. This method of ranking does assume approximately equal yields of scFv from each clone. Since this is not always the case, some scFvs may also be identified that simply express high levels of scFv. These can be discriminated from those of higher affinity by further characterization of the scFvs (see Example 18).

Preparation of ScFv from 1ml E.coli Cultures

[0741] Individual E.coli colonies containing a phagemid representing one of the unique scFvs from panel 3 were inoculated into 96-well plates containing 100 μl 2TYAG medium per well. Eight wells on each plate were reserved for positive and negative control samples. The plate was grown overnight at 30°C with shaking at 120 rpm.

Next day, 1ml of 2TYAG + 345 mM sucrose was added to each well of an autoclaved 96 deep well plate (Beckman). Twenty μ l of each overnight culture was resuspended and transferred to the appropriate well of the deep well plate. The plate was grown for approximately 3.5 hours at 30°C with shaking at 250 rpm (or until the OD₆₀₀ = 0.6). Fifty μ l of 1M IPTG was added to 5ml 2TY and 10 μ l of this was added to each well. The plate was grown overnight at 30°C with shaking at 250 rpm.

Plates were kept at 4°C for the remainder of the procedure. The overnight plate (above) was centrifuged at 3500 rpm for 10 minutes at 4°C to pellet the cells. The supernatant was decanted and each pellet resuspended in 100µl TES (0.2M Tris HCl pH8.0, 0.5mM EDTA, 0.5M sucrose) and transferred to a fresh 96 well plate. This plate was incubated on ice for 30 minutes and then centrifuged for 10 minutes at 3500 rpm at 4°C to pellet the cell debris. During centrifugation, 15µl of freshly made protease inhibitors cocktail (Roche, 1 tablet dissolved in 1.5 ml water) was added to each well of a fresh 96 well plate. Supernatants from the centrifuged plate were then transferred to the plate containing the protease inhibitors. The plate was centrifuged at 3500 rpm for 10 minutes at 4°C and the supernatant was transferred to a further 96-well plate. This step was repeated at least once more or until there was no sign of any cell debris following

centrifugation. Finally, the plate was covered in foil to prevent evaporation of samples during the BIAcore run.

Generation of a high density HIS-BLyS surface

[0744] All BIAcore analysis was performed on BIAcore 2000 machines, using the BIAcore 2000 control software and evaluated using the BIAevaluation 3.0 software. A high density His-tagged BLyS surface (>1000 RU HIS-BLyS coupled) was prepared in flow cell 2 by amine coupling to a CM5 chip. A new CM5 chip was inserted into the BIAcore and a sensorgram started over flow cell 2 with HBS buffer at a flow rate of 5μl/min. The NHS and EDC solution were mixed 1:1 before injecting 30μl over the CM5 surface. Fifty μl HIS-BLyS (at 10μg/ml in Sodium acetate buffer, pH4) was injected and allowed to couple to the surface. Thirty μl of ethanolamine-HCl solution was then injected to block free NHS esters. Prior to using the chip, 10μl of 4M Guanidine hydrochloride in HBS was injected over the surface to strip the surface of non-covalently bound BLyS. A blank surface (no HIS-BLyS) was also prepared over flow cell 1 so that non-specific binding effects can be subtracted from the HIS-BLyS binding curves.

[0745] Typically, a 5000 RU His-tagged BLyS surface was generated in this way and used for 96-well analysis of scFvs isolated from the periplasm of E.coli.

BIAcore Analysis

[0746] The 96-well plate containing periplasmic scFvs was secured inside the BIAcore. Two ml of 4M Guanidine hydrochloride in HBS was placed in a rack inside the BIAcore for regeneration of the HIS-BLyS surface between samples. The sensorgram was run over flow cells 1 and 2 at a flow rate of 20µl/minute. The following method was run:

MAIN

FLOWCELL 1,2,3,4

LOOP cycle STEP
APROG inj %pos
ENDLOOP

APPEND CONTINUE

END

DEFINE LOOP cycle

LPARAM %pos

rla1

rlb1

rlc1

r1d1

rlel

rlfl etc (all wells listed until rlh12)

END

DEFINE APROG inj

PARAM %pos

FLOW 20

KINJECT %pos 35 30 !scfv injection

QUICKINJECT r2f3 10 !regeneration

EXTRACLEAN

END

[001] When the run had finished, the sensorgram data for flow cell 1 was subtracted from the data for flow cell 2 for each sample using the BIAevaluation software. The clones were compared with one another principally by overall RU change as the scFv dissociates from the surface. In addition a few scFvs were identified as having potentially slower off-rates. An example of the dissociation section of a typical sensorgram for 8 scFvs is shown in Figure 14. An anti-TNFα antibody that does not recognize BLyS was included as a control. Of the 8 scFvs exemplified, I079F06 was identified for further study due to the relatively high numbers of RU's bound to the surface.

[0748] ScFvs were identified principally if they demonstrated a RU change of over 1200, a few were also identified as having potentially slower than typical off-rates. A total

of 28 clones were chosen on these criteria and are listed in Table 7.

Table 7: Identification of 28 antibodies to membrane-bound BLyS that demonstrate the most significant RU changes by BIAcore

Antibody	Antibody
I079C01	I084C04
I082H08	I080E05
I079E02	I083B12
I079B05	I082G01
I079F06	I082G02
I079F08	I082C03
I079F11	I082A05
I079B12	I082D07
I080B01	I082B08
I080G09	I084A01
I099D03	I084B02
I080D03	1080A08
I080A03	I084C11
I083G03	
I080G07	

Example 18: scFv Affinity Determinations

[0749] The affinity (K_D) of the 28 scFvs was determined using the BIAcore.

Low Density HIS-BLyS Surface for Kinetic Studies

[0750] 500RU surfaces were used for kinetic studies of purified scFv binding to HIS-BLyS. The method to prepare these surfaces was identical to the method described in Example 17, only smaller volumes of HIS-BLyS were injected.

Measurement of scFv Binding Kinetics

[0751] The chip containing the low density HIS-BLyS surface was inserted into the BIAcore. A dilution series for each of the 28 purified scFvs (prepared as in Example 6) were diluted in HBS (typically starting with 50µg/ml scFv and double diluting down to 1.5µg/ml). The dilution series was then injected sequentially over the blank control (flow cell 1) and low density HIS-BLyS surface (flow cell 2) using the following program:

MAIN

FLOWCELL 1,2,3,4

APROG	genab	rld1	ab1
APROG	genab	r1d2	ab2
APROG	genab	rld3	ab3
APROG	genab	rld4	ab4
APROG	genab	r1d5	ab5
APROG	genab	r1d6	ab6

APPEND CONTINUE

END

DEFINE APROG genab

PARAM %Abpos %AbId

FLOW 20

KINJECT %Abpos 200 80

INJECT r2f3 10

EXTRACLEAN

END

[0752] Bound scFv were removed by injecting 10µl of 4M Guanidine hydrochloride in HBS (location r2f3 in the above program) over the surface between samples. Binding curves for individual scFv were analysed using the BIAevaluation software to determine antibody on- and off-rates.

[0753] A typical example of the binding curves generated for the scFv antibody I082C03 is shown in Figure 15. The off-rate for this clone was calculated as $2x10^{-3}$ s⁻¹. The affinity of I082C03 was calculated as 20 nM, assuming 100% activity of the scFv. The 5 scFvs with the highest affinities as scFvs are given in Table 8.

Table 8: Identification of 5 antibodies to membrane-bound BLyS that have the highest affinities as scFvs

Antibody	Affinity
	(K _D)
I079F11	5nM
I079E02	10nM
I082G02	6nM .
I082H08	. lnM
I099D03	4nM

Example 19: Recognition of mouse membrane-bound BLyS

[0754] The ability of the 5 scFvs listed in Table 8 to also recognize murine membrane-bound BLyS was determined using a competition ELISA. This assay assesses the ability of test phage scFvs to bind to the membrane-bound form of BLyS on the murine cell line, P388, plasma membranes in the presence of different forms of competing human BLyS. Competing BLyS was either presented as the His-tagged form of BLyS, or soluble BLyS. ScFvs that recognize mouse membrane-bound BLyS would give an ELISA signal on the P388 plasma membranes that is competed out by pre-incubation with HIS-tagged BLyS but not by pre-incubation with soluble BLyS.

Maintenance of P388.D1 cells and preparation of plasma membranes

[0755] P388.D1 cells are a mouse monocyte-macrophage like cell line. They were cultured in L-15 medium supplemented with 2mM L-glutamine, 10% CS, 10U penicillin, 100g/ml streptomycin (all reagents from Sigma). Cells were split 1:4 every 3-4 days to maintain a cell density of 2-8 x 10⁵ per ml. A fresh aliquot of cells was thawed from liquid nitrogen every 6 weeks. Plasma membrane fractions were prepared as described in Example 2.

Competition ELISA

[0756] P388 plasma membranes (50µl per well) were coated at 10µg/ml in PBS onto Falcon 96-well plates overnight at 4°C. The method is otherwise essentially as described Example 16.

[0757] The results for 3 clones, I079E02, I082H08 and I099D03 are shown in Figure 16. All 3 scFvs recognize P388 plasma membranes. This binding is competed out in the presence of HIS-tagged BLyS (HIS-BLyS) but not in the presence of biotinylated BLyS (bio-BLyS). This confirms that these scFvs also recognize the membrane-bound

form but not the soluble form of mouse BLyS.

Example 20: Conversion of scFvs to IgG1 format

[0758] The VH domain and the VL domains of scFvs that we wished to convert into IgG molecules were cloned into vectors containing the nucleotide sequences of the appropriate heavy (human IgG1) or light chain (human kappa or human lambda) constant regions such that a complete heavy or light chain molecule could be expressed from these vectors when transfected into an appropriate host cell. Further, when cloned heavy and light chains are both expressed in one cell line (from either one or two vectors), they can assemble into a complete functional antibody molecule that is secreted into the cell culture medium. Methods for converting scFvs into conventional antibody molecules are well known within the art.

Generation of NSO cell lines expressing anti-BLyS antibodies (IgG1)

[0759] Plasmids containing the heavy and light chains were separately linearized using the Pvu I restriction enzyme. The linearized DNAs were purified by phenolchloroform extraction followed by ethanol precipitation and then resuspended in H₂O. NS0 cells (10⁷) from a growing culture were electroporated (0.25kV and 975µF) in PBS with 12.5 µg linearized heavy chain plasmid DNA and 37.5 µg linearized light chain DNA. The cells were washed in 20 ml non-selective medium (10% FCS in DMEM supplemented with 6mM glutamine, amino acids and penicillin/streptomycin) and then transferred in 12.5 ml medium into a T75cm² flask and incubated overnight at 37°C, 5% CO2/air. The day after transfection the cells were resuspended in selective medium containing 1mg/ml geneticin and dispensed into 5 x 96-well plates at 200 µl/well. After 18 days at 37°C (5% CO₂/air) the colony supernatants were screened by an ELISA that detects assembled human IgG in order to identify colonies expressing IgG. Approximately twenty positive colonies were expanded and adapted to growth in serumfree, selective medium. Duplicate T25cm² flasks were set up. Cells from one flask were frozen down as a stock and cells in the second flask were grown to saturation. The productivity of the saturated cultures was assessed by ELISA. The highest producing cell lines were then selected for large-scale antibody production.

[0760] The above procedure is exemplified for the I006D08 anti-BLyS antibody

constructs. Following electroporation and selection of NSO cells, supernatants from ninety-three wells each containing a single colony were screened by ELISA to detect assembled IgG1, antibody. Twenty-seven of the supernatants were identified as containing IgG. The colonies from 24 of the positive wells were transferred to 1ml selective medium in a 24-well plate and allowed to grow for 2 days. The 1ml cultures of cells were then added to 4ml selective medium containing reduced serum (0.5% FCS) in a T25cm² flask. When the cultures reached confluency 1 ml cells were diluted in 4ml selective, serum-free medium in a T25cm² flask. At confluency this subculture regime was repeated again. Finally 1ml cells from the culture containing 0.1% FCS was diluted with 9 ml serum-free, selective medium and divided into 2 x T25cm² to form the saturated and stock cultures. The stock cultures were frozen down and stored in liquid nitrogen once the cultures were confluent. The saturation culture was grown until the viability of the culture was < 10%. Twenty-three out of the 24 colonies originally expanded were successfully adapted to growth in serum-free medium. The productivity of these serum-free adapted cell lines ranged from 0.3 to 17 µg/ml by ELISA quantification of the saturated, 5ml serum-free cultures. The I006D08-32 cell line produced 17 μg/ml.

Large-scale IgG production

[001] The highest-producing cell lines were revived from frozen stocks and then expanded to 400ml in selective, serum-free medium in 2 liter roller bottles. The cells were grown at 37°C and rolled at 4 rpm with the headspace being re-equilibrated with 5% CO₂/air every 2-3 days. Finally the culture was expanded to a 4 liter volume by the addition of serum-free medium without selection (400 ml per 2 liter roller bottle). The cultures were then grown to saturation.

[001] This procedure is exemplified by the production of I006D08 antibody from the I006D08-32 cell line. The frozen stock of I006D08-32 was revived into a T25 cm² containing 5 ml serum-free medium containing 1mg/ml geneticin and grown at 37°C in 5% CO₂/air incubator. After two days growth the culture was diluted with 7.5 ml fresh medium and transferred to a T75cm² flask. After a further three days in the incubator the cells were transferred to 130 ml selective medium and transferred to a 2 liter roller bottle. After three days growth the cells were diluted with 500 ml selective medium and split into 2 x 2 liter roller bottles. After another 2 days 100 ml fresh selective medium was added to

each roller. Finally the next day the culture was expanded to a total volume of 4 liters with non-selective medium and divided into 10×2 liter roller bottles. After three days the medium was supplemented with 6mM glutamine. The cells were grown for 17 days from the final subculture into a 4 liter volume. The cells grew up to 3×10^6 cells/ml before viability declined to $< 0.2 \times 10^6$ cells/ml. At this low viability the culture supernatants were harvested. ELISA analysis indicated that the culture supernatant contained 33 μ g/ml IgG. Hence, the 4 liter culture contained 132 mg IgG.

IgG Purification

[0763] The purification of the IgG from the fermentation broth is performed using a combination of conventional techniques commonly used for antibody production. Typically the culture harvest is clarified to remove cells and cellular debris prior to starting the purification scheme. This would normally be achieved using either centrifugation or filtration of the harvest. Following clarification, the antibody would typically be captured and significantly purified using affinity chromatography on Protein A Sepharose. The antibody is bound to Protein A Sepharose at basic pH and, following washing of the matrix, is eluted by a reduction of the pH. Further purification of the antibody is then achieved by gel filtration. As well as removing components with different molecular weights from the antibody this step can also be used to buffer exchange into the desired final formulation buffer.

Purification of I006D08 IgG1

[001] The harvest was clarified by sequential filtration through 0.5 μm and 0.22 μm filters. Clarified harvest was then applied to a column of recombinant Protein A Sepharose equilibrated at pH 8.0 and washed with the equilibration buffer. I006D08 antibody was eluted from the Protein A Sepharose by application of a buffer at pH3.5. The collected antibody containing eluate was then neutralized to pH 7.4 by the addition of pH 8.0 buffer. The neutralized eluate was concentrated by ultrafiltration using a 30 KDa cut off membrane. Concentrated material was then purified by Sephacryl S300HR gel filtration using phosphate buffered saline as the mobile phase. The final monomeric IgG1 fraction from the gel filtration column was then concentrated to the desired formulation

concentration by ultrafiltration using a 30 KDa cut off membrane. The final product was filtered through a 0.22 μm filter.

Example 21: Antibody neutralization of murine splenocyte proliferation as measured by 3HdT incorporation

[0765] To determine if an antibody inhibited BLyS mediated B cell proliferation, a splenocyte proliferation assay was performed Briefly, murine splenocytes were isolated by flushing spleen with complete medium using a 25g needle and 10 ml of complete medium (RPMI 1640 with 10% FBS containing 100U/ml penicillin, $100\mu g/ml$ streptomycin, 4mM glutamine, 5×10^{-5} M β -mercaptoethanol). The cells were passed through a 100 micron nylon filter to remove cell clumps. The cell suspension was then ficolled at 400 x g for 25 minutes at room temperature (one 15 ml conical tube/spleen; 3 ml ficol, 10 ml cell suspension/spleen; Ficol 1083 from Sigma). The recovered cells were washed 3 times in complete medium and counted. Recovered cells were then diluted to a concentration of 3×10^6 /ml in complete medium containing a 3X concentration of SAC (3X = 1:33,333 dilution of stock) (Staph. aureus Cowan strain; Calbiochem).

[0766] For each antibody, 50 microliters of antibody dilutions at 30µg/ml, 3.0µg/ml, and 0.3µg/ml concentrations were aliquotted into individual wells of a 96 well plate in triplicate. Suitable positive controls, such as, for example monoclonal antibody 15C10, were also used. Medium containing no antibody (and human isotype controls (purchased commercially) when necessary) were used as negative controls.

[0767] BLyS protein was diluted in complete medium to concentrations of 300ng/ml, 90ng/ml and 30ng/ml. 50 microliters of each of the BLyS dilutions were then added to the antibody dilution series in the plates. The plate containing the antibody and BLyS dilutions are then incubated for 30 minutes at 37°C, 5% CO₂, after which 50 microliters of the splenocyte cell suspension containing SAC was added to all wells. The plates were then incubated for 72 hours (37°C, 5% CO₂).

[0768] After 72 hours, each well was supplemented with 50µl of complete medium containing 0.5µCi of 3H-thymidine (6.7 Ci/mM; Amersham) and cells were incubated for an additional 20-24 hours at (37°C, 5% CO₂). Following incubation cells were harvested using a Tomtec Cell Harvester and filters counted in a TopCount

Scintillation counter (Packard).

Example 22: Human B cell proliferation assay for in vitro screening of BLyS antagonist molecules

[0769] The bioassay for assessing the effects of putative BLyS antagonists was performed in triplicate in 96 well format by mixing equal volumes of BLyS, responder cells, and putative antagonist each of which is prepared as a 3X stock reagent.

[0770] B-lymphocytes were purified from human tonsil by MACS (anti-CD3 depletion), washed, and resuspended in complete medium (CM) (RPMI 1640 with 10% FBS containing 100U/ml penicillin, 100µg/ml streptomycin, 4mM glutamine, 5x10E-5 M beta-mercaptoethanol) at a concentration of 3 x 10e6 cells/mL. Staphylococcus aureus, Cowan I (SAC, CalBiochem) was added to cells at 3X concentration (3X = 1:33,333 dilution of stock

[0771] Meanwhile, eight serial dilutions (3-fold) of potential antagonist were prepared in CM such that the diluted antagonists are at 3X the final concentrations to be tested in the assay. Antibodies are routinely tested starting at a final concentration of 10ug/mL and going down to about 1.5 ng/mL.

[0772] Human rBLyS was prepared in CM to 3X concentration (3X = 300 ng/mL, 30 ng/mL, and 3 ng/mL) in CM. Potential inhibitors were routinely tested at several concentrations of BLyS to avoid false negatives due to unexpectedly low affinity or antagonist concentration.

[0773] Fifty microliters of diluted antagonist and 50uL of diluted BLyS were added to the putative antagonist dilution series.

[0774] Cells were then incubated for 72 hours (37°C, 5% CO₂) in a fully humidified chamber. After 72 hrs., the cells were supplemented with 0.5 µCi/well 3H-thymidine (6.7 Ci/mmol) and incubated for an additional 24 hours. Plates were harvested using a Tomtec Cell Harvester and filters counted in a TopCount Scintillation counter (Packard).

The entire disclosure of each document cited (including patents, patent applications, journal articles, abstracts, laboratory manuals, books, or other disclosures) in this application is incorporated in their entireties herein by reference. Additionally, the specifications and sequence listings of U.S. Provisional Applications Nos. 60/212,210 filed June 16, 2000; 60/240,816 filed October 17, 2000; 60/276,248 filed March 16, 2001; 60/277,379 filed March 21, 2001; and 60/293,499 filed May 25, 2001 are all hereby incorporated by reference in their entireties.

Table 1: scFvs that Immunospecifically Bind to BLyS

Table 1: scFvs that Immunospecifically Bind to BLyS

			•		VO: 2132)	_			2131)																					
VH CDR3 Sequence (SEQ ID NO)		HDDDVLTGYYFES (SEQ ID NO: 2130)	SRDLLLFPHYGMDV (SEQ ID NO: 2133)	DRYDILTGYYYYGMDV (SEQ ID NO: 2129)	VQMDSEYYDLLTGINVGPYYFDY (SEQ ID NO: 2132)	DGYYDILTGYSYYGMDV (SEQ ID NO: 2135)	GYDSSAFRAFDI (SEQ ID NO: 2136)	APYDLLTHYFHYFDY (SEQ ID NO: 2134)	AATTSQKHNKYAYYFYGMDV (SEQ ID NO:	PFYDTL TSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQVWVA (SEQ ID NO: 2143)	PFYDTLTSYVFQVWVA (SEQ ID NO: 2143)	PFYDTLTRYVFQYFDH (SEQ ID NO: 2144)	PFYDTLTGYVFQYFDH (SEQ ID NO: 2141)	PFYDTLTRYVFQVWVA (SEQ ID NO: 2142)	PFYDTLTGYVFQVWVA (SEQ ID NO: 2140)	PFYDTLTRYVFQYFDH (SEQ ID NO: 2144)	PFYDTLTGYVFQYFDH (SEQ ID NO: 2141)	PFYDTLTRYVFQVWVA (SEQ ID NO: 2142)	PFYDTLTRYVFQVWVA (SEQ ID NO: 2142)	PFYDTLTGYVFQVWVA (SEQ ID NO: 2140)	PFYDTLTGYVFQVWVA (SEQ ID NO: 2140)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQVWVA (SEQ ID NO: 2143)	PFYDTLTSYVFQVWVA (SEQ ID NO: 2143)	PFYDTLTRYVFQYFDH (SEQ ID NO: 2144)	PFYDTLTRYVFQYFDH (SEQ ID NO: 2144)	PFYDILTSYVFQYFDH (SEQ ID NO: 2139)	PFYDILTSYVFQYFDH (SEQ ID NO: 2139)
AAs of	VH CDRS	99-111	99-112	102 - 117	99 - 121	99 - 115	99 - 110	99 - 113	99 - 118	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114		99 - 114	99 - 114	$\overline{}$	7	7		,	99 - 114
AAs of	CDR2	50 - 66	20 - 66	52 - 69	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	99 - 05	99-05	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66		20 - 66	20 - 66	1.		20 - 66	20 - 66	20 - 66	20 - 66
AAs of	CDR1	26-35	26-35	26-37	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	.26-35	26 - 35	26-35	26 - 35	26-35	26 - 35	26-35	26 - 35	26-35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	t	26 - 35
AAs of	ΗΛ	1-122	1 - 123	-1 - 128	1 - 132	1 - 126	1-121	1 - 124	1-129	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125
AAs of	CDR3	228 - 237	228 - 238	234 - 243	234 - 244	228 - 238	228 - 240	229 - 239	236 - 245	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240		232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	i.	•	232 - 240
AAs of	CDR2	189 - 195	189 - 195	195 - 201	195 - 201	189 - 195	189 - 195	190 - 196	197 - 203	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	-		•	193 - 199
AAs of	CDRI	160 - 173	163 - 173	166 - 179	169 - 179	163 - 173	160 - 173	164 - 174	168 - 181	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177
AAs of	۸۲	138 - 248	139 - 249	144 - 254	148 - 255	142 - 249	137 - 251	140 - 250	145 - 256	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251	141 - 251
scFv	SEC E	1		m	4	S	9	7	∞	<u>.</u>	10	11	17	13	14	15.	16	. 11	18	19	20	77	22	73	74	. 25	5 6	. 12	78	23
Clone ID		1003F12S	1006D08	I008A11	I017D10	1022D01	I031F02	1050A12	I051C04	1050B11	I050B11-01	I050B11-02	I050B11-03	I050B11-04	I050B11-05	I050B11-06	I050B11-07	I050B11-08	I050B11-09	1050B11-10	1050B11-11	I050B11-12	I050B11-13	I050B11-14	I050B11-15	I050B11-16	I050B11-17	I050B11-18	I050B11-19	1050B11-20

ON O	PRYDII TRYVEOVEDH (SEC ID NO: 2138)	(SEC ID NO.		C Z	į	įŻ	Ċ	PFYDTL TSYVFOYFDH (SEO ID NO: 2137)	PFYDTLTSYVFOVWVA (SEO ID NO: 2143)	PFYDILTSYVFOYFDH (SEO ID NO: 2139)	PFYDTLTSYVFOYFDH (SEO ID NO: 2137)	PFYDTLTSYVFOYFDH (SEO ID NO: 2137)	Ö	Ö	CN CL		PFYDTLTSYVCRPHF (SEO ID NO: 2238)	(SEO ID NO	(SEO ID	(SEO ID NO	SEO ID NO.	(SEO ID NO:	(SEQ ID NO:	PFYDTLTSYVPCSPPR (SEQ ID NO: 2261)	PRYDTLTSYVCYPPA (SEQ ID NO: 2240)	PFYDTLTSYVLPLLS (SEQ ID NO: 2224)	PFYDTLTSYVALYRL (SEO ID NO: 2234)	PFYDTLTSYVRASFS (SEO ID NO: 2271)	PFYDTLTSYVCTPVP (SEQ ID NO: 2319)	PFYDTLTSYVWPSFFS (SEQ ID NO: 2277)	PFYDTLTSYVTPRGY (SEQ ID NO: 2275)	PFYDTLTSYVSSLLS (SEQ ID NO: 2213)	PFYDTLTSYVPLLPLC (SEQ ID NO: 2263)	PFYDTLTSYVPPPSFL (SEQ ID NO: 2266)
99 - 114	99 - 114					99 - 114	99 - 114	99 - 114		99 - 114	99 - 114	99 - 114	99 - 114	99 - 114		99 - 114	•	99 - 113	99 - 113	99 - 113	99-113	99 - 113	99 - 113	99-114	99 - 113	99 - 113	99 - 113	99-113	99-113	99 - 114	99 - 113		99 - 114	99 - 114
50 - 66	50 - 66	50-66	50 - 66	•	50 - 66	50 - 66	20 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	99 - 09	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66		•	1	-1	20 - 66	20 - 66
26 - 35		26 - 35	26 - 35	26 - 35	26 - 35		26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35		26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35			26 - 35		26 - 35	26 - 35					26 - 35	26 - 35
1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1-125	1 - 125	1 - 124	1 - 124	1 - 124	1 - 124	1 - 124	1 - 124	1 - 124	1 - 125	1 - 124	1 - 124	1 - 124	1 - 124	1 - 124	1 - 125	1 - 124	1 - 124	1 - 125	1 - 125
232 - 240	•	•	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	231 - 239	231 - 239	•	231 - 239	231 - 239	231 - 239	•	•	•	ı	•	٠	٠,	•	•	•	1	232 - 240
3 - 199		3 - 199	3 - 199	3 - 199	3 - 199	3 - 199	3 - 199	3-199	3 - 199	3 - 199				3 - 199	3-199	3 - 199	2 - 198	192 - 198	•					•				•	•			٠		- 199
7 193 7 193	• •	7 193	7 193	7 193	7 193			7 193	7 193	Ξ	_			7 193	7 193	7 193						_						_		_	_		_	7 193
166 - 177 166 - 177	- 1	166 - 177	166 - 177	166 - 177	166 - 177	166 - 177	7	166 - 177	166 - 177	166 - 177	$\overline{}$		166 - 177	166 - 17	166 - 177	166 - 177	•		•		<u> </u>		•				1			1	•	165 - 176	166 - 177	166 - 177
-251 -251	•	-251	-251	-251	-251	-251	- 251	-251	-251	•	٠		•	-251	-251	-251	- 250	- 250				- 250	- 250	- 251	- 250			•	•		. 1		- 251	-251
141	141	141	141	141	141	141	141	141	. 141	141	141	141	141	14	141	141	140-	140-	140	140	140	140	140 -	141	140-	4	140	140	9	141	9 :	3 :	141	141
30	32	33	34	35	36	37	38	33	40	41	42	43	4	45	46	47	48	49	20	51	25	23	24	55	26	57	28	20	9	[9	3 5	3 3	94	S
I050B11-21 I050B11-22	I050B11-23	I050B11-24	I050B11-25	1050B11-26	I050B11-27	I050B11-28	· 1093D03	I093D09	I093G08	I097D11	II01A04	1101B01	I102A02	1102E01	1102G06	I087A07	I087A08	I087A09	1087B02	I087B03	I087B04	1087B05	1087B06	1087B08	1087809	1087C02	1087C05	1087C06	I087C07	1087C08	1087D01	108/D02	108/D03	108/1005

																	•									
PFYDTLTSYVPTSTT (SEQ ID NO: 2269) PFYDTLTSYVISCSWA (SEQ ID NO: 2299) PFYDTLTSYVSALPPP (SEQ ID NO: 2274) PFYDTLTSYVCRH F (SEQ ID NO: 2274)	PFYDTLTSYVVSFPSL (SEQ ID NO: 2307) PFYDTLTSYVMGVTPS (SEQ ID NO: 2322)	FFYDILISYVLFRPVL (SEQ ID NO: 2326) PFYDILISYVPSVGG (SEQ ID NO: 2267)	PFYDTLTSYVPPTRH (SEQ ID NO: 2286) PFYDTI TSYVI BORD (SEQ ID NO: 2243)	PFYDTLTSYVPLLPP (SEQ ID NO:	PFYDTLTSYVLRCVL (SEQ ID NO: 2239) PEYDTT TEXYTHEEDS (SEQ ID NO: 2239)	PEYDILISYVLRLPPO (SEO ID NO. 2263)	PFYDTLTSYVGPYGT (SEQ ID NO: 2284)	PFYDTLTSYVTTPCT (SEQ ID NO: 2276)	ASYLSTSSSLDN (SEQ ID NO: 2265)	PEVDTI TSYVIPEI PI (SEQ ID NO: 2137)	PFYDTLTSYVLHIYPH (SEO ID NO: 2235)	PFYDTLTNYVFEYYAS (SEO ID NO: 2323)	PFYDTLTSYVILYYLH (SEQ ID NO: 2295)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVLMYFPH (SEQ ID NO: 2220)	PEYDILISYVLFFYPL (SEQ ID NO: 2325)	PEYDTLTSYVFDYYAS (SEO ID NO: 2137)	PFYDTLTSYVIPELPL (SEQ ID NO: 2290)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFEYYSL (SEQ ID NO: 2324)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVLEFYLL (SEQ ID NO: 2303)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVLPLDS (SEQ ID NO: 2223)	PFYDTLTSYVLYFYPS (SEQ ID NO: 2317)
99 - 113 99 - 114 99 - 114	1 1	1 1	99 - 113		99 - 113				99-110	99 - 114	99 - 114	99 - 114			7:	99 - 114			99 - 114	99 - 114		•	1			99 - 114
50 - 66 50 - 66 50 - 66 50 - 66	50-66	20-00	50 - 66 50 - 66		50 - 66 50 - 66	50-66	50 - 66	50 - 66	20 - 66	50 - 66	50 - 66	99-09	20 - 66	20 - 66	90-00	20-00	50 - 66	50 - 66	•	•	•				20-66	20 - 66
26-35 26-35 26-35 26-35	1 1		26-35		26-35		26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35		•		1	•		26-35	20-32	26 - 35
1 - 124 1 - 125 1 - 125 1 - 124	1-125	1 - 124	1 - 124	1 - 124	1-124	1-125	1 - 124	1-124	1 - 121	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1-125	1 - 125	1 - 125	1 - 125	1 - 125	1-125	1 - 125	1-125	1 - 124	1 - 125
231 - 239 232 - 240 232 - 240 231 - 239	232 - 240	1 1	231 - 239 231 - 239		231 - 239 231 - 239	•	4	•	225 - 233		- 1	1	1	•	087 - 787					•	•	•	•	•	•	737 - 740
192 - 198 193 - 199 193 - 199 192 - 198	193 - 199 193 - 199 193 - 199	• •	192 - 198 192 - 198	•	192 - 198 192 - 198				186 - 192		193 - 199				193 - 199					•		١.	•	•	•	193 - 199
165 - 176 1 166 - 177 1 166 - 177 1 165 - 176 1	166 - 177 1 166 - 177 1 166 - 177 1	176	65 - 176 L	- 176	65 - 176 19 65 - 176 19	-177	-176	- 176	00 - 1/0 13 66 - 177 16	- 177	-177	- 177	- 12	13	- 177	. 177	-171	171	-177	-177	- 177	- 177	- 1771	- 177	9/1-	00-1// 13
140 - 250 1 141 - 251 1 141 - 251 1 140 - 250 1	141 - 251 1 141 - 251 1 141 - 251 1	-250	140 - 250 1 140 - 250 1		140 - 250 1 140 - 250 1			140 - 250 I	15/-244 1	•	- 251 1	-251 1	-251 1	141-251 1	- 251	-251	- 251	-251	-251	- 251	-251 1	167-	1 107 -	141 - 251 1	1 052-	1 107-
66 68 69	852	5 25 5	4 5	92		79	& .	≅ 8															•			
1087D07 1087D09 1087E04 1087E05	I087E10 I087F02 I087F04	1087F05	108/F0/ 1087F08	1087F09	1087G06 1087G06	I087G07	1087G09	108/G10	108/F102 1088A01	I088A03	1088A04	I088A08	1088A09	1088A10	1088A12	1088B01	I088B02	I088B03	1088B05	1088B06.	1088BU/	TOSETOS	10881509	1088B10 1088B10	100801	700007

PFYDTLTSYVFQYFDH (SEQ ID NO: 2137) PFYDTLTSYVLHYYAL (SEQ ID NO: 2215) PFYDTLTSYVLHYYAL (SEQ ID NO: 2225) PFYDTLTSYVFQYFDH (SEQ ID NO: 2137) PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)		(SEQ ID NO: (SEQ I	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)
99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114			
50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66			4-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1
26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35			
1 - 125 1 - 125 1 - 125 1 - 125 1 - 125 1 - 124 1 - 125	1-125 1-125 1-125 1-125 1-125 1-125 1-125	1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125	1 - 125 1 - 125 1 - 125 1 - 125 1 - 125 1 - 125 1 - 125
232 - 240 232 - 240 232 - 240 232 - 240 232 - 240 231 - 239 232 - 240	232 - 232 -	232- 232- 232- 232- 232- 232- 232-	232 - 232 -
193 - 199 193 - 199 193 - 199 193 - 199 193 - 199 193 - 199			1 1 1 1 1 1 1 1
166 - 177 166 - 177 166 - 177 166 - 177 165 - 176 165 - 176		' 	166 - 177 166 - 177 166 - 177 166 - 177 166 - 177 166 - 177
141 - 251 141 - 251 141 - 251 141 - 251 140 - 250 141 - 251		141 - 251 141 - 251	
102 103 104 105 106 108	5 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	122 123 124 128 128	130 131 132 134 135 136
1088C03 1088C09 1088C12 1088D01 1088D03 1088D04 1088D04	1088D11 1088D11 1088E01 1088E02 1088E03 1088E04 1088E10 1088E11	1088G03 1088G03 1088G07 1088G10 1088H05 1092A03 1092A06 1092A06	1092A10 1092A11 1092B01 1092B02 1092B04 1092B10 1092B10

											٠																							
PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVLALDL (SEQ ID NO: 2328)	ÿ:	PRYDIT TOWN WWITH (SEQ ID NO: 2137)	PRYDIT TSYVEOVEDH (SEQ ID NO: 2228)	(SEO ID NO	SEO ID NO.	SEC ID NO.	(SEO ID NO:	(SEO ID NO:	SEO ID NO	(SEO ID NO	SEO ID NO:	(SEO ID NO	(SEO ID NO	(SEO ID NO	(SEO ID NO:	(SEO ID NO:	(SEO ID NO:	(SEO ID NO:	(SEO ID NO:	(SEO ID NO:	(SEO ID NO:	(SEQ ID NO:	D NO: 2265)	PFYDTLTSYVLPVYDH (SEQ ID NO: 2334)	PFYDTLTSYVFQYFAH (SEQ ID NO: 2268)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	D NO:	Ö	PFYDILISYVIFYYPT (SEQ ID NO: 2289)	PFYDTL TSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVLEVYHP (SEQ ID NO: 2318)	ö	
99 - 114	•	99 - 114		i .	99 - 114	•			99 - 114	99 - 114	99 - 114	99 - 114	99 - 114		99 - 114		99 - 114				99 - 114	99 - 114	99 - 114	99 - 110		7	99-114		99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	
50 - 66		20-06			50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	50 - 66	50 - 66	99 - 09	50 - 66	20 - 66	99 - 09	99 - 09	50 - 66		20 - 66		•	•		20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	
26 - 35		26 - 35 26 - 35			.3	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35		26 - 35	í	•	1	•	•		•	٠	26 - 35	
1 - 125 1 - 125	1 - 124	1 - 125			1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1-121	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	I - 125	1 - 125	
232 - 240 232 - 240		232 - 240			232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240		•	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	•		•	•	•	•	•			•			232 - 240	32 - 240	
193 - 199 :	192 - 198	8 2	- 199	- 199	- 199	93 - 199	93 - 199	- 199	- 199	- 199	- 199	- 199	- 199	- 199	- 199	- 199	- 199	199	- 199	- 199	- 199	- 199	- 199	- 192	- 199	- 199	- 199	- 199	- 199	- 199	- 199	- 199	3-199 2	
-177 -	65 - 176 19	.177	- 177	- 177	- 177	- 177 1	-177	-177 1	-171	- 177	- 177	- 177	- 177	-177	- 177	-177	- 177	- 177	-177	- 177	- 177	- 177	-177	- 170	-177			_ `					- 177 193	
251 1 251 1	250 1	251 1	_	_	<u> </u>	251 166	≃ _	9	–	=	Ξ.	–	9	9	9	Ξ.	~	=	19	9	19	9	251 166	9	251 166	901 107				_ `			251 166	
	140	141 -	141-	. 141 -	t	•	•	٠		•	€.	•	•	٠	•		•	•	٠	•	•		141 -		•		•		٠	1	•	•	141 -2	
138 139	140	142	143	14	145	146	147	148	149	120	151	152	153	154	155	156	157	158	159	9	191	162	<u> </u>	104	165	001	797	80.	69	2:	Z :	7/1	173	
1092C01 1092C02	1092C07	1092C12	1092D01	I092D07	1092D09	I092D10	I092D11	1092E01	1092E03	1092E04	1092E07	1092E10	1092E11	I092F01	1092F02	I092F05	I092F07	1092F08	I092F11	I092F12	1092G01	1092G05	1092610	1092H01	1093A06 1003 A 00	1093A09	1093A11	1093A12 1003D02	1093B02	1093805	1093506	1093509	1093B12	

PFYDTLTSYVLHAYAF (SEQ ID NO: 2332) PFYDTLTSYVFQYFDH (SEQ ID NO: 2137) PFYDTLTSYVILYYLH (SEQ ID NO: 2295)	FFYDILISY VFEFLPL (SEQ ID NO: 2245) PFYDTLTSYVRPFYAH (SEQ ID NO: 2273) PFYDTLTSYVFOYFDH (SEO ID NO: 2137)	PFYDTLTSYVLHFYRV (SEQ ID NO. 2302)	PFYDTLTSYVIQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVHEFFSL (SEQ ID NO: 2283) PFYDTI TSYVMOEEPT (SEQ ID NO: 2221)	PFYDTLTSYVLSFYPV (SEQ ID NO: 2246)	PFYDTLTSYVLYYYAF (SEQ ID NO: 2251)	FFYDILISY VFQYFDH (SEQ ID NO: 2137) PFYDTLTSYVFOYFDH (SEQ ID NO: 2137)	S C C C C C C C C C C C C C C C C C C C	PFYDTLTSYVLHFYPL (SEQ ID NO: 2333)	BNO	D NO:	Ö	(SEQ ID NO:	PEYDTI TEXAT BYWAYS (SEQ ID NO: 2280)	(SEO ID NO:	(SEO ID	D NO:	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	9	Ö G G		PEYDTLTSYVEOVEDH (SEQ ID NO: 2137)	D NO	(SEQ ID NO:	PFYDTLTSYVLEAFSL (SEQ ID NO: 2311)	PFYDTLTSYVFGFYPF (SEQ ID NO: 2252)
99 - 114 99 - 114 99 - 114				99 - 114			99 - 114		_			99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114				99 - 114			99 - 114	99 - 114	99 - 114
50 - 66 50 - 66 50 - 66		1 1		50-66	50 - 66	20 - 66	50 - 66 50 - 66	20 - 66	99 - 09	20 - 66	50 - 66	20 - 66	20 - 66	20 - 00 50 - 66	50-66	50 - 66	20 - 66	20 - 66		00-00			50 - 66	50 - 66	20 - 66	20 - 66
26 - 35 26 - 35 26 - 35 26 - 35		26 - 35		26 - 35	٠,		26 - 35							26 - 35	26-35	26-35	26 - 35	26 - 35		26 - 35	26 - 02	26 - 35		26 - 35	26 - 35	26 - 35
1 - 125 1 - 125 1 - 125 1 - 125	1 - 125 1 - 125 1 - 125	1 - 125	1 - 125	1-125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1-125	1-125	1 - 125	1 - 125	1 - 125	1-125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125
232 - 240 232 - 240 232 - 240 232 - 240		232 - 240		232 - 240 232 - 240		•	232 - 240 232 - 240			•	•		1	232 - 240		232 - 240		•	232 - 240	047 - 757				1	232 - 240	232 - 240
193 - 199 193 - 199 193 - 199		193 - 199		193 - 199 193 - 199		193 - 199	- 199	- 199	- 199	66!	- 199	- 199	266		-199	- 199	- 199		193 - 199	100	100	18	193 - 199	- 199	- 199	193 - 199
66 - 177 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	-177	66 - 177 1	171	.66 - 177 1 .66 - 177 1	177	66 - 177 1	111	171	177	177	177	- 177	111	-177	-117	-177	- 177	- 177	1 1/1 1	. 177	177	-177	66-177 1	- 177		6-177
-251 16 -251 16 -251 16		-251 16 -251 16	. —	-251 16 -251 16	_	-251 16		_	_ :	_ :			01 167-	-					-251 16	-	-	251	-251 16	_	_	-251 16
4 141 S 141 6 141		141		4 4 14 14 14		141							141						141				5 141			141
174 175 176 177	178	181	18.	183	18,	¥ 6	188	189	<u>ج</u>	191	192	193	194	196	197	198	19	5 00	707	203	200	205	200	207	208	70
1093C02 1093C03 1093C05 1093D05	1093D08 1093D10	1093D12 1093E01	1093E02	1093E03 1093E08	1093E10	1093F01 1002F02	1093F05	1093F08	1093F11	1093G07	1093G11	1063017	1007.27.00	1094B07	I094B08	1094B12	1094C11	1094C12	10941706	1094D08	10941009	1094D10	1094D11	1094E04	1094E08	1094F04

	PFYDILISYVEQYEDH PFYDALTSYVEQYEDH PFYDTLTSYVEQYEDH
	99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114 99 - 114
50 - 66 50 - 6	50 - 66 50 - 66
	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35
1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125	1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125 1-125
	232 - 240 232 - 240
199 199 199 199 199 199 199 199	193 - 199 193 - 199
6 - 177 1 1 1 1 1 1 1 1 1	66 - 177 1 1 1 66 - 177 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
-251 16-2	251 1 251 1
210 141 211 141 213 141 214 141 219 141 219 141 222 14	
444444444444444444444444444444444444444	
1094F05 1094F11 1094F11 1094F12 1094G06 1095A04 1095B04 1095B09 1095C02 1095C09 1095C09 1095C09 1095C09 1095C09 1095C09	1095D09 1095E01 1095E05 1095F06 1095G06 1095G01 1096A10 1096B01 1096B01 1096C00 1096C00 1096C00

																		-																		
	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137) PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	(SEQ ID NO:	(SEQ ID NO:	(SEQ ID NO:	(SEQ ID NO:	ö	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVIHELPL (SEQ ID NO: 2330)	PFYDTLTSYVIPFLPL (SEQ ID NO: 2290)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVMHYLPV (SEQ ID NO: 2257)	PFYDTLTSYVLEFFSH (SEQ ID NO: 2315)	ID NO: 2265)	PFYDTLTSYVIHYLVT (SEQ ID NO: 2294)	E NO	SEQ1	(SEQ	(SEQ ID NO:	ON CI	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	(SEQ ID NO:	(SEQ ID NO		PFYDTLTSYVIHFYSL (SEQ ID NO:	PFYDTLTSYVFGFFPH (SEQ ID NO:	PFYDTLTSYVFQYFDH (SEQ ID NO:		ASYLSTSSSLDN (SEQ ID NO: 2265)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)					
	99 - 114 99 - 114		n'	•		1			1		99 - 114	99 - 114	•				1		•			1	1						99 - 114	•					99 - 114	
	50 - 66 50 - 66	99 - 09				٠	•				20 - 66	20 - 66	20 - 66		20 - 66	1	- 1		20 - 66	20 - 66	20 - 66		20 - 66	1	•			20 - 66	20 - 66	20 - 66	20 - 66	•	20 - 66	•	20 - 66	
	26 - 35 26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	t	26 - 35	26 - 35	26 - 35	•	•		26 - 35	26 - 35	26 - 35		٠		,		က	•	က္		26 - 35	26 - 35	26 - 35		26 - 35	26-35	26 - 35	
	1 - 125 1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 121	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 121	1 - 125	٠
	232 - 240 232 - 240	•	•	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	t	225 - 233	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240		232 - 240	•	232 - 240	232 - 240	232 - 240	232 - 240	232 - 240	•		232 - 240	
	193 - 199 193 - 199	93 - 199	193 - 199	1	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	186 - 192	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	7	7	ï	ï	193 - 199	193 - 199	193 - 199	193 - 199	193 - 199		186 - 192	193 - 199	
	66 - 177 1	- 177	-177	-177	66 - 177 1	66 - 177	66-177]	66 - 177 1	66 - 177		- 177	66 - 177	66 - 177	66 - 177	60 - 170	66 - 177	66 - 177	66 - 177	66-177	- 177	- 177	66 - 177	66 - 177	66 - 177	66 - 177	66-177	. 66 - 177	66-177	120 - 177	66 - 177	721 - 991	166 - 177	120 - 177	60 - 170	120 - 177	
-	141 - 251 1 141 - 251 1		141-251	141-251	141 - 251	141-251 1	41-251	141 - 251 1	41-251 1	41-251 1	41-251 1	41 - 251 1	41 - 251 1	41 - 251	37 - 244 1	41 - 251	41 - 251 1	41 - 251 1	41-251 1	41 - 251	[41 - 25]	41-251	41 - 251	41 - 251	41-251	141 - 251	141 - 251	141 - 251	41 - 251	41 - 251	41 - 251	41 - 251		37 - 244	141 - 251	
	246 1 247 1		_		,	_				7	_	_	_	_	_		_	_		_	_	_	-	_	_	_		•;•		276	_	_		280	•	
	1096D01 1096D02	1096D05	. 90О9601	1096D09	I096E02	1096E06	1096E11	I096F02	1096G01	1096G02	1096G05	1096G07	60D960I	I096G12	10H960I	1097A04	1097A06	I097A09	I097B02	1097B09	I097B10	I097B11	1097C05	1097C09	I097C11	1097D05	1097D06	1097E01	1097E04	1097E08	I097E09	I097F09	1097G10	I097H02	I098A04	

																		•																
PFYDTLTSYVFQYFDH (SEQ ID NO:	PLYDILISY VLDFYSV (SEQ ID NO:	PEVILLISI VEQIFUH (SEQ ID NOT)	PEVILLISI VLI I IAF (SEQ ID NO:	DEVINIT TEXT REFERENCE (SEC ID NO.	DEVICE TEXTED VEFF IFF (SECTION NO.)	PRYTHE TENTED VED (SEQ ID NO:	PEVDET TO THE CASE OF THE SECTION OF	PENDIT TOWN FOREDHI (SEQ ID NO:	PEVILLISI VEQUEDII (SEQ ID NO:		PRVDTI TSVVI BEEDD (SEQ ID NO.	PFYDTI.TSYVEOVEDH (SEO ID	PEYDTL TSYVEOVEDR (SEO ID NO:	PFYDTI.TSYVEOYEDH (SEO ID NO:	PFYDTLTSYVFOYFDH (SEO ID NO:	PFYDTLTSYVFOYFDH (SFO ID	PFYDTLTSYVFOYFDH (SEO ID	PFYDTLTSYVFOYFDH (SEO ID	PFYDTLTSYVFOYFDH (SEO ID	PFYDTLTSYVFOYFDH (SEO ID NO:	PFYDTLTSYVLHYYAH (SEO ID NO.	PFYDTLTSYVFQYFDH (SEQ ID NO:	PFYDTLTSYVFQYFDH (SEQ ID NO:	PFYDTLTSYVFQYFDH (SEQ		PFYDTLTSYVFQYFDH (SEQ	•	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQYFDH (SEQ ID NO:	PFYDTLTSYVFQYFDH (SEQ ID NO:	D NO:	(SEQ ID NO:	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)	PFYDTLTSYVFQYFDH (SEQ ID NO: 2137)
	99 - 114					1	1							•				99 - 114		99 - 114	99 - 114	99 - 114				•			99 - 114	99 - 114	7	7	99 - 114	99 - 114
50 - 66	5 6	50.		•					20-05	50 - 66			50 - 66	50 - 66		50 - 66	20 - 66	50 - 66	50 - 66	99 - 09	20 - 66	20 - 66	20 - 66		20 - 66	20 - 66	20 - 66		•	20 - 66	•		ŧ	20 - 66
26 - 35	26-										. m	۳,	26 - 35	26 - 35	.3	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	ر ا	er i	•	. ن			. ع	٠,		•		26 - 35	26 - 35
1 - 125	1 - 125	1 - 125		1 - 125	1 - 125			1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	51-1	1-125	1-125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1-125	1-125
232 - 240		•			•	•	ŧ			•			232 - 240	232 - 240	232 - 240	232 - 240		•	•	•	•	•	1	ı	•	•		•	t	•	•	232 - 240	232 - 240	732 - 240
193 - 199	-		193 - 199	193 - 199	193 - 199	193 - 199	193 - 199	7	7	193 - 199	193 - 199	193 - 199	93 - 199	193 - 199			•		- 199	- 199	- 199	- 199	- 199	2 2	7 2	5	661	561	- 199	56]	661 -	961 -	•	. 661 - 6
166 - 177	- 177	- 177	- 177	_	66-177 1	166-177 1	_	- 177	166 - 177 1	166 - 177 1	166 - 177 1	-177	- 177	- 177	-177	- 177	- 177	- 177	- 177	- 177	- 177	- 177		1771		1 //1 -		//[-	///-	//!-	. 771 -	1	1771	00-1/1
-251	-251	-251	-251	-251	-251 1	-251	-251	- 251	-251	-251	-251	-251	- 251	-251	-251	-251	-251	-251	-251	-251	-251	- 251	167-	991 167-	_							01 167 -		_
282 141 283 141				87 141	88 141	89 141	90 141	11 141	2 141	3 141						_	-						141	_	_	7 -						141		•
N 14	284	7	Ä	7	7	7	73	53	23	23	22	χ,	, <u>22</u>	Χ ί	%	, <u>2</u> ,	∺ ;	∺ :	₩ 8	<u> </u>	,	35	א ה	2 2	2 6	3 2	7 5	10.	2 5	ָר. ק		10.	7 .	;
1098A05 1098B08	I098C01	1098C04	I098F11	I098F12	I098G02	I098G12	1098H05	1101A01	1101B04	1101B06	1101D04	1101D07	1101E09	1101E12	1101G02	1101611	1102C03	1102E09	1102F02	1102508	1102400	1106B02	1106B04	1106007	1106E05	1106R12	11066011	1106001	1100000	110011	1109011	1109G012	1109H04	1000000

೯೫೯			•
PFYDTLTSYVFQYFDH (SEQ ID NO: 2137) PFYDTLTSYGFQYFDH (SEQ ID NO: 2232) PFYDTLTSYVFQYFDH (SEQ ID NO: 2137) SRDLLLFPHHGLDS (SEQ ID NO: 2146) SRDLLLFPHHSFDL (SEQ ID NO: 2147) SRDLLLFPHHGLDY (SEQ ID NO: 2151) SRDLLLFPHHGLDV (SEQ ID NO: 2151) SRDLLLFPHHSLDL (SEO ID NO: 2152)	SRDLLLFPHHSFDL (SEQ ID NO: 2147) SRDLLLFPHHALSP (SEQ ID NO: 2147) SRYLLLFPHHSFDL (SEQ ID NO: 2149) SRVLLLFPHHSFDL (SEQ ID NO: 2147) SRDLLLFPHHSFDL (SEQ ID NO: 2477) SRDLLLFPHDHLLF (SEQ ID NO: 2639) SRDLLLFPTHPLSF (SEQ ID NO: 2561) SRDLLLFPTHPLSF (SEQ ID NO: 2550) SRDLLLFPLAPLFF (SEQ ID NO: 2550) SRDLLLFPSDPLSL (SEQ ID NO: 2559)	(SEQ ID SEQ ID S	SRDLLLFPLSPLSF (SEQ ID NO: 2574) SRDLLLFPDFPMAP (SEQ ID NO: 2433) SRDLLLFPHSPLY (SEQ ID NO: 2470) SRDLLLFPQDPLSP (SEQ ID NO: 2372) SRDLLLFPDDPLLS (SEQ ID NO: 2430) SRDLLLFPGPLLI (SEQ ID NO: 2400) SRDLLLFPHGPLLI (SEQ ID NO: 2491) SRDLLLFPGSPLLF (SEQ ID NO: 2491) SRDLLLFPTAALSF (SEQ ID NO: 2341) SRDLLLFPTAALSF (SEQ ID NO: 2341)
PFY PFY SRD SRD SRD SRD SRD SRD SRD SRD SRD	SRU	S C C C C C C C C C C C C C C C C C C C	S S S S S S S S S S S S S S S S S S S
99 - 114 99 - 114 99 - 115 99 - 112 99 - 112 99 - 112			99 - 112 99 - 112 99 - 111 99 - 112 99 - 112 99 - 112
8888888	999999999999999999999999999999999999999	99999999999	888888888
8 8 8 8 8 8	50.	50-10-20-20-20-20-20-20-20-20-20-20-20-20-20	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8
26 - 35 26 - 35		26 - 35 26 - 3	26 - 35 26 - 35
1 - 125 1 - 125 1 - 125 1 - 123 1 - 123 1 - 123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123
232 - 240 232 - 240 232 - 240 228 - 238 228 - 238 228 - 238 228 - 238			228 - 238 228 - 238 227 - 237 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238
199 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2			2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
193 - 1 193 - 1 193 - 1 189 - 1 189 - 1 189 - 1			189 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -
######################################			2222222
166 - 1 166 - 1 163 - 1 163 - 1 163 - 1 163 - 1			163-1 162-1 163-1 163-1 163-1 163-1 163-1 163-1
251 251 251 249 249 249 249			249 249 249 249 249
141 141 139 139 139		139 - 249 139 - 249 138 - 248 139 - 249 139 - 249 139 - 249 139 - 249 139 - 249	139 - 139 -
318 320 321 322 323 324 324	326 327 328 329 330 331 332 333	335 336 337 338 340 342 343	345 346 347 348 350 351 353
1110B03 1112D09 1112F10 1089F12 1105E12 1108E06 1113E07	1114G05 1116A01 1116A09 1116C11 1085A01 1085A03 1085A04 1085A05	1085A06 1085A07 1085A09 1085A10 1085B01 1085B02 1085B03 1085B04 1085B04	1085B06 1085B07 1085B10 1085B12 1085C02 1085C03 1085C03 1085C05

PCT/US01/19110

EQ D NO: 2 SEQ D NO: 2 SEQ D NO: 2 SEQ D NO: 3 SEQ D	SRDLLLFPAAPLLF (SEQ ID NO: 2403)
22 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	99 - 112
	20 - 66
26 - 35 26 - 3	26 - 35
	1 - 123
227 - 237 228 - 238 228 - 238	228 - 238
194 195 195 195 195 195 195 195 195 195 195	189 - 195
1	163 - 173
249 249 249 249 249 249 249 249 249 249	139 - 249
354 354 355 356 357 357 358 358 358 358 358 358 358 358 358 377 378 378 378 378 378 378 378 378 37	389
1085C09 1085C10 1085C10 1085D01 1085D03 1085D04 1085D04 1085D03 1085D03 1085D11 1085E01 1085E01 1085E01 1085E01 1085E01 1085F02 1085F03 1085F03 1085F04 1085F04 1085F04 1085F06 1085F06 1085F07 1085F07 1085F07 1085F07 1085F07 1085F07 1085F07 1085F07 1085F07 1085F07	1085G03

2135)	:		. '
SRDLLLFPSAPLDP (SEQ ID NO: 2601) SRDLLLFPNAVLDI (SEQ ID NO: 2629) SRDLLLFPSEPLFF (SEQ ID NO: 2664) SRDLLLFPSSVLWP (SEQ ID NO: 2338) SRDLLFPHAPLQ (SEQ ID NO: 2354) SRDLLFPDSPLAP (SEQ ID NO: 2445) SRDLLLFPSSPLHP (SEQ ID NO: 2445) DGYYDILTGYSYYGMDV (SEQ ID NO: 2135) SRDLLLFPSMPLTF (SEQ ID NO: 2695) SRDLLLFPHSILHP (SEQ ID NO: 2438) SRDLLLFPHSILHP (SEQ ID NO: 2438) SRDLLLFPHAPI SH (SEQ ID NO: 2569)	SRDLLLFPDAALRF (SEQ ID NO: 222) SRDLLLFPSSHLSF (SEQ ID NO: 2421) SRDLLLFPSAPLSS (SEQ ID NO: 2624) SRDLLLFPHAPLTP (SEQ ID NO: 2577) SRDLLLFPHFPLHF (SEQ ID NO: 2335) SRDLLLFPHFPLHF (SEQ ID NO: 2348) SRDLLLFPHFPLHF (SEQ ID NO: 2348)	SRDLLLFPEPLII (SEQ ID NO: 2457) SRDLLLFPASPLNP (SEQ ID NO: 2364) SRDLLLFPSSPLYF (SEQ ID NO: 2720) SRDLLLFPTSPLSF (SEQ ID NO: 2579) SRDLLLFPDDGLSS (SEQ ID NO: 2428) SRDLLLFPISPLCF (SEQ ID NO: 2530) SRDLLLFPTAPLYG (SEQ ID NO: 2535) SRDLLLFPHHSLFF (SEQ ID NO: 2427) SRDLLLFPQGPLRF (SEQ ID NO: 2427) SRDLLLFPQGPLRF (SEQ ID NO: 2440) SRDLLLFPQGPLRF (SEQ ID NO: 2440)	SEQ ID NO: SEQ ID NO: SEQ ID NO: SEQ ID NO: SEQ ID NO: (SEQ ID NO: SEQ ID NO: (SEQ ID NO:
99 - 112 99 - 112 99 - 112 99 - 112 99 - 112 99 - 115 99 - 115		99 - 112 99 - 112	99-112 99-112 99-112 99-112 99-112
50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35
1 - 123 1 - 123	1-123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1-123 1-123 1-123 1-123 1-123 1-123
228 - 238 228 - 238 228 - 238 228 - 238 227 - 237 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238		228 - 238 228 - 238	228 - 238 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238
195 195 195 195 195 195 195 195	198 198 198 198 198 198 198 198 198 198	195 195 195 195 195 195	195 195 195 195 195
173 189 173 189 173 189 172 188 173 189 173 189 173 189 173 189		173 189 173 189 173 189 173 189 173 189 173 189 173 189 173 189 173 189	173 189 173 189 173 189 173 189 173 189 173 189
		163	
139 - 249 139 - 249		139 - 249 139 - 249 139 - 249 139 - 249 139 - 249 139 - 249 139 - 249	139 - 249 139 - 249 139 - 249 139 - 249 139 - 249 139 - 249
390 391 392 394 397 398 398	402 403 404 405 406 407	408 409 410 411 411 411 411 411 411 411 411 411	418 420 421 423 424 424
1085G04 1085G07 1085G08 1085G10 1085G11 1085G12 1085H10 1086A03 1086A04	1086A07 1086A09 1086A10 1086A11 1086A12 1086B02	1086B05 1086B06 1086B07 1086B09 1086B11 1086C03 1086C05 1086C05	1086C09 1086C10 1086C11 1086D01 1086D04 1086D05

2365)	33	2665)	<u>ج</u>	2610)	2469)	(17	(8)	<u> </u>	86 8	Ş:	€(3 6	3 =	5	5 6	<u> </u>	2 6	26		÷6	∵ (<u> </u>	२	<u>ر</u>	- 6	2	_ ′	<u>_</u> (و	<u></u>	ر ا	<u>ල</u>	<u> </u>	ବ୍ରଚ	•
Ö	Ö	ö	ġ Ş	ÿ	D.NO: 24	NO: 2621)	NO: 2598	(SEQ ID NO: 2567)	(SEQ ID NO: 2398)	(SEC ID NO: 2490)	(SEQ 1D NO: 2464)	(SEO ID NO: 2507)	SRDLLLFPSSPLRI (SFO ID NO: 2714)	NO. 2540)	(SEO ID NO: 2642)	(SEO ID NO: 2652)	(SEO ID NO: 2033)	(SEO ID NO: 2513)	SRDI I FPFDPI I (SEO ID NO: 2454)	(950 ID NO. 2434)	NO. 233	NO: 2407	NO: 2448)	NO: 2383)	NO: 2591)	107 : ON	7: 2012)	0.2390)	(SEQ ID NO: 2483	NO: 2539)		NO: 2436)	NO: 25/2)	NO: 2450) NO: 2147)	1
(SEQ ID NO	(SEQ ID	(SEQ ID	a :			SEQ ID NO:					SEC ID NO.	SEC E	EO EO	SRDLI, FPTEPI OF (SEO ID NO:	A CHO		A CHI	SEO E		, EO 19	36					SRDI I I FEHDEI I (SEO ID MO: 26)		SPDI I EPHTEI UE (CEO ID NO.		(SEQ ID NO:	۹.		∃ 6		
			SLEF (ALFIE ALOS			_ ,		DI DE	PIRE	PLRI (S	PLOF	DI CA	. `	PITE	PI DI	PI 1176	AT DI		מונס				3) 11 TO	30,171	מבותה עם							
SRDLLLFPHTHLTF	SRDLLLFPHSSLDF	SKULLFFNHPMFP	OKULLEPLOSLEF (LLFFIN	SECULLIFIER AFILIES	SEDITIFIED ALOR	יימייי די	SKULLEFRIFLIF (SKULLFFAAHLSF SPDI I I EDD A DI I E	SPDI I TERRETLE	SP DI LI EDP A DI NE	SRDIJI FPTAPI RE	LEFPSS	L. FPTF	SRDLL FPSDPI SA	SRDLLLEPYNPPIF	SRDI.I. FPHTPI I E	SRDI I.I FPHAPI DI	I FPFD	SRDLL FPTDAT PT	SPINIT TERMANT	SPDI I I EDBCDI I D	SRDI I I FPV A PI CE	SPILL I FDANCI CE	SRDI I I FPVCPI TU	I EDUT	1 507 1	SPDI I I FRUTTRI UE	111111	SKULLEF LUALYF	SKULLEPY I FLLF	SKULLLFFHUPLIF	SEDELEFFE I LUF	SRDLLLFPHHSFDL	
SRDL	SRDL	SKUL	TO SE	מים בי	לים המים	מל בי		CDU	CPUL	200		SRDL	SRDL	SRDL	SRDL	SRDLI	SRDI	SRDLI	SRDL	SRDL	SPDI	SPDIT	SRDIT	וועמא	SEDIT	SPDIT	L Lugo	SPDI		SPEC		SECUL	משטו ז	SRDLL	
•		211 - 66	•			7	7	7 7	7 7				99 - 112	99 - 112	99 - 112				•	Ţ				٠,				7	•			00 - 112	7 7	7	
9;	8 %	_				3 %	3 4		-		۔ ورو	99	.99	99	99	92	. 99	9	. 99	99	عر.	3 12					_							8 9	
	, ,	•	•				8 6						50-	- 09	- 92	50-	50 -	50-		50 -		20	20.	20-	50 - (50-		50-1			•	. 1			
	26 - 35											•	26 - 35	26 - 35	26-35	26 - 35	26-35	26-35	26 - 35	26 - 35	٠,	יא נ	. "	. (1	. 6	(7)	"	י אי	י ר		26 - 36			i m	
1 - 123	1 - 123	1 - 123	1 - 123		1 - 123		1-123			1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 122	1 - 123	1 - 123	1 1 1 3	123	123	- 123	- 123	1- 123	
-238	238	238	- 238	- 238	238	- 238	. 238	.238	. 238	238	. 238	. 238	. 238	. 238	.238	238	238	238	238	238	238	238	238	238	238							238	238 1	238	
228				•			228		•	228.	228	228	228 -	228 -	228 -	228 -	228 -	228 -	228 -	228 -	228 -	228	228 -	228 -	228 -	227 -	228 -	228 -	228 -			228	228	228 -	
89 - 195				9 - 195		9-195	9 - 195	•		9-195	9-195		•		- 195	1	- 195	9-195	- 195	- 195	- 195	- 195	- 195	- 195	- 195	- 194	- 195	- 195	- 195						
173 18	73 18	-	_	173 189	73 189	73 189	173 189	173 189	73 189	173 189	3 189	_	_	73 189		73 189	3 189	3 189	3 189	3 189	3 189	3 189	3 189	3 189	3 189	2 188	3 189	3 189	3 189	_	_	_		_	
163 - 17		7	7	163 - 17	163 - 17	163 - 17	163 - 17	163 - 17	163 - 17	163 - 17	163 - 173		7	_	7	-	163 - 17	163 - 17	163 - 17	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	62 - 17	163 - 173	163 - 173	63 - 173	•	63 - 173	7	63 - 173	63 - 173	
139 - 249	-249	139 - 249	- 249	- 249	- 249	139 - 249	- 249	- 249	- 249	139 - 249	- 249	- 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	249	139 - 249		249	249	248	249	249	249	249	249	- 249]	- 249 1	- 249	
139	139	139	139	139	139	139	139 - 24	139 - 24	139	139	139	139	139	139	139	139	139	139	139	139.	139	139	139	139	139 - 24	138-	139 -	139 -	139 -	139	139 -	139-	139 -	139 -	
426	428	429	430	431	432	433	434	435	436	437	438	439	9 3	441	442	443	4	445	446	447	448	449	450	451	452	453	454	455	456	457	458	459	460	461	
D07	D09	D10	DII	D12	E02	E03	E05	E07	E08	E09	E10	E12	F02	٠ د د	8 0.	603	F11	303	304	305	906	307	906	310	105	101	103	901	. 201	801	110	111	301	. 20	
1086D07 1086D08	1086D09	I086D10	I086D11	I086D12	I086E02	I086E03	I086E05	1086E07	1086E08	I086E09	1086E10	1086E12	1086F02	1086F05	1086F08	1086F09	1086F11	1086G03	1086G04	1086G05	90D980I	1086G07	1086G09	I086G10	1086H05	I089A0	I089A03	1089A06	I089A07	1089A08	I089A10	I089A11	1089B01	I089B02	

SRDLLLFPTSPLQP (SEQ ID NO: 2528)	SRDLLLFPSSPI.IF (SEO ID NO: 2712)			_	SRDLLLFPKSPILF (SEO ID NO: 3	•						SRDLLLFPSPYLSF (SEQ ID NO: 2701)	SRDLLLFPQAPLFD (SEQ ID NO: 2683)	SRDLLLFPHAPFTF (SEQ ID NO: 2507)	SRDLLLFPHAPLVL (SEQ ID NO: 2581)	SRDLLLFPHYGMDV (SEQ ID NO: 2133)	SRDLLLFPHYPLLF (SEQ ID NO: 2344)	SRDLLLFPSSPLSP (SEQ ID NO: 2717)	SRDLLFPHAPLFT (SEQ ID NO: 2546)	SRDLLLFPNDPLLI (SEQ ID NO: 2634)	SRDLLLFPHAPLQ (SEQ ID NO: 2554)	SRDLLLFPSHAFHE (SEQ ID NO: 2677)	SRDLLLFPNHPLYP (SEQ ID NO: 2663)	SRDLLLFPYSPLFP (SEQ ID NO: 2657)	SRDLLLFPQDPLHP (SEQ ID NO: 2346)	SRDLLLFPDAPLFP (SEQ ID NO: 2423)	SRDLLLFPHSPLLI (SEQ ID NO: 2453)	SRDLLLFPGSPLLF (SEQ ID NO: 2491)	SRDLLLFPSSPLTF (SEQ ID NO: 2718)	SRDLLLFPTQPLSF (SEQ ID NO: 2566)		SRDLLLFPTFPLLF (SEQ ID NO: 2380)	(SEQ ID NO	SRDLLLFPYSPLLF (SEQ ID NO: 2670)
99 - 112		- 1	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99-112	99 - 112		•				ì			•	99 - 112	99 - 112		99 - 112	ŧ	99 - 112
99 - 09	20 - 00 50 - 66		99 - 09	20 - 66	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	99 - 05	99 - 09		20 - 66	20 - 66	99 - 09	99-05	20 - 66 	20 - 66	99 - 09		20 - 66		•	•	•	•	•		•	•		•	ı	20 - 66
26 - 35	26 - 35			26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35			26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	•							•	•	1					26 - 35	26 - 35
1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123		1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 122	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123
228 - 238			228 - 238	228 - 238	228 - 238	228 - 238	228 - 238			•	228 - 238				•	•	228 - 238	228 - 238	•		•	•	•		١.	•	•	٠	•		•	•		228 - 238
189 - 195	ı (189 - 195	189 - 195	189 - 195	189 - 195		189 - 195	٠		1						•	189 - 195	189 - 195	189 - 195			189 - 195	189 - 195	189 - 195	189 - 195	•		٠	•	•			•	189 - 195
163 - 173		163 - 173	163 - 173	163 - 173	163 - 173	1	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	1	1	•	ï	163 - 173	163 - 173			•		4	•		•	•				163 - 173	163 - 173	•	163 - 173
139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	138 - 248	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	•		139 - 249
462	464	465	466	467	468	469	470	471	472	473	474	475	476	477	478	479	480	481	482	483	484	482	486	487	488	489	490	491	492	493	494	495	496	497
1089B03 1089B04	1089B05	1089B06	I089B07	I089B08	I089B09	I089B10	I089B11	1089C01	1089C02	I089C03	1089C05	1089C06	1089C07	1089C09	I089D01	1089D02	I089D03	1089D04	I089D05	I089D07	1089D08	1089D09	I089D11	I089E01	1089E02	1089E03	1089E04	1089E06	1089E09	1089E10	1089E11	1089F01	1089F03	10891-04

	2135)	
SRDLLLFPHSPLRI (SEQ ID NO: 2459) SRDLLLFPRAPLIF (SEQ ID NO: 2490) SRDLLLFPLAPLSF (SEQ ID NO: 2567) SRDLLLFPROPLSF (SEQ ID NO: 2565) SRDLLLFPROPLSF (SEQ ID NO: 2565) SRDLLLFPSAPLTF (SEQ ID NO: 2565) SRDLLLFPSAPLTF (SEQ ID NO: 2687) SRDLLLFPSAPLTF (SEQ ID NO: 2721) SRDLLLFPGSPLTF (SEQ ID NO: 2389) SRDLLLFPTAPLLF (SEQ ID NO: 2389) SRDLLLFPAPLLF (SEQ ID NO: 2389) SRDLLLFPAPLLF (SEQ ID NO: 2567) SRDLLLFPSAPLDF (SEQ ID NO: 2567)	2671) D NO: 2416) 2678) 2678) 2678) 2648) 2648) 2600) 2709)	(SEQ ID NO: SEQ ID NO:
1 - 123 1 - 123	1 - 123 1 - 123	1 - 123 1 - 123
- 195 - 195 - 195 - 195 - 195 - 195 - 195 - 195 - 195 - 195	195 195 195 195 195 195 195 195 195 195	195 195 195 195 195 195 195 195
		173 173 173 173 173 173 173 173
249 249 249 249 249 249 249 249 249 249	249 249 249 249 249 249	249 16 249 16 249 16 249 16 249 16 249 16 249 16
498 499 500 501 503 504 505 506 508 509 510		524 525 526 527 529 529 530 531
1089F05 1089F06 1089F08 1089F10 1089F11 1089G01 1089G03 1089G05 1089G05 1089G05	1089G11 1089H10 1090A02 1090A03 1090A04 1090A05 1090A07 1090B01 1090B01	1090B05 1090B05 1090B06 1090B11 1090C01 1090C03 1090C05 1090C06

		•									
1090C08	534	139 - 249	163 - 173	3 189.	195	228 - 238	1 - 123	26 - 35	99 - 09	99 - 112	SRDLLLFPYSPLSF (SEO ID NO: 2676)
1090C10	535	139 - 249	163 - 17	3 189.	. 195	228 - 238	1 - 123	26 - 35	99-09	99 - 112	ID NO:
I090D02	536	139 - 249	163 - 173	3 189.	. 195	228 - 238	1 - 123	26 - 35	99-09	99 - 112	D NO:
I090D03	537	139 - 249	163 - 173	3 189.	. 195	228 - 238	1 - 123	26 - 35	99 - 09	99 - 112	D NO:
1090D04	538	139 - 249	163 -	_	. 195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	ID NO:
I090D05	539	139 - 249	163 -	3 189-	195	228 - 238	1 - 123	26 - 35	30 - 66	99 - 112	SRDLLLFPHAPLFE (SEQ ID NO: 2529)
1090D06	240	139 - 249	163 -	_	. 195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPRAPLDF (SEQ ID NO: 2367)
1090D07	241	139 - 249	163 -]		. 195		1 - 123	•	99 - 09	99-112	SRDLLLFPFGTLRF (SEQ ID NO: 2462)
1090D08	542	139 - 249	_	3 189.	. 195	228 - 238	1 - 123	26 - 35	50 - 66	99 - 112	D NO:
1090D09	543	139 - 249	163 -	3 189-	195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPRDPLAF (SEQ ID NO: 2505)
1090D12	54	139 - 249		3 189-	. 195	228 - 238	1 - 123	26 - 35	99 - 09	99 - 112	SRDLLLFPTSPLSF (SEQ ID NO: 2579)
1090E04	545	139 - 249	163 -	3 189-	195	228 - 238	1 - 123	26 - 35	99 - 09	99 - 112	SRDLLLFPHAPLLL (SEQ ID NO: 2552)
1090E05	246	139 - 249		3 189	195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPSAPISF (SEQ ID NO: 2588)
1090E06	547	139 - 249	_	3 189.	195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPQGPLSF (SEQ ID NO: 2443)
I090E07	548	139 - 249		3 189.	195	228 - 238	1 - 123	26 - 35	50 - 66	99 - 112	SRDLLLFPGSPLHP (SEQ ID NO: 2484)
1090E09	549	139 - 249	163 -	3 189	. 195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPSDPLSF (SEQ ID NO: 2647)
1090E11	250	139 - 249	163 -	_	195	228 - 238	1 - 123	•	20 - 66	99 - 112	SRDLLLFPHDGLAP (SEQ ID NO: 2700)
I090E12	551	139 - 249	163 -	189	- 195		1 - 123	•		99 - 112	SRDLLLFPTSPLTF (SEQ ID NO: 2582)
1090F01	552	139 - 249		3 189.	195	228 - 238	1 - 123	26 - 35	99 - 09	99 - 112	SRDLLLFPNGPLHP (SEQ ID NO: 2649)
I090F02	553	139 - 249	_	3 189.	195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPQAPLSF (SEQ ID NO: 2696)
I090F03	554	139 - 249	-	189	- 195	228 - 238	1 - 123	26 - 35	50 - 66	99 - 112	
1090F04	555	139 - 249	_	189	- 195	228 - 238	1 - 123	26 - 35	99 - 09	99 - 112	SRDLLLFPFFPLQF (SEQ ID NO: 2460)
I090F05	256	139 - 249	163 -	189	- 195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPLDPLHF (SEQ ID NO: 2359)
1090F06	557	139 - 249	163 -	_	195	228 - 238	1 - 123	26 - 35	99 - 0 9	99 - 112	D NO:
I090F07	258	139 - 249	163 -		195	.0	1 - 123	٠	20 - 66	99 - 112	SRDLLLFPFAPLRF (SEQ ID NO: 2451)
1090F08	559	139 - 249		_	195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPLHPLIF (SEQ ID NO: 2570)
1090F09	260	139 - 249	163 -	_	195		1 - 123			•	SRDLLLFPHYPLLF (SEQ ID NO: 2344)
I090F10	561	139 - 249	163 -		195		1 - 123		•		
I090F11	262	139 - 249	163 -	_	195	•	1 - 123	۳,	20 - 66	•	SRDLLLFPSNPLTF (SEQ ID NO: 2698)
1090G01	2 63	139 - 249	163 -	_	. 195		1 - 123	1	20 - 66	99 - 112	SRDLLLFPTAPLEI (SEQ ID NO: 2347)
1090G02	564	139 - 249		_	. 195	٠	1 - 123	26 - 35	•	99 - 112	SRDLLLFPRDPLQF (SEQ ID NO: 2395)
1090G04	265	139 - 249	_	3 189-	. 195	228 - 238	1 - 123	26-35		1	SRDLLLFPHEPLAF (SEQ ID NO: 2633)
1090G05	266	7	163 -	_	195	228 - 238	1 - 123	26 - 35	20 - 66	•	
1090G06	267	139 - 249	_	_	195	228 - 238	1 - 123	26 - 35	20 - 66	1	(SEQ
1090G07	268	139 - 249	163 - 17	3 189-	195	1	1 - 123	ر ع	20 - 66		(SEQ
1090G08	869	139 - 249	163 - 17	3 189.	. 195	228 - 238	1 - 123	26 - 35	20 - 66	99 - 112	SRDLLLFPHTPLTF (SEQ ID NO: 2492)

SRDLLLFPSEPLRI (SEQ ID NO: 2356) SRDLLLFPTAPLDF (SEQ ID NO: 2343) SRDLLLFPNRGLDL (SEQ ID NO: 2669) SRDLLLFPYDPLFM (SEQ ID NO: 2724) SRDLLLFPSAPLAF (SEQ ID NO: 2592) SRDLLLFPSAPLAF (SEQ ID NO: 2594) SRDLLLFPHSPITF (SEQ ID NO: 2441) SRDLLLFPRYPLFF (SEQ ID NO: 2481)	(SEQ ID NO. (SEQ I	(SEQ ID NO: (SEQ I
	*	
8 6 8 8 8 8 8	222222222222222222222222222222222222222	******
50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 6	50 - 66 66 66 66 66 66 66 66 66 66 66 66 6
26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 3	26 - 35 2 26 - 3
1-123 1-123 1-123 1-123 1-123 1-123 1-123	1-123 1-123 1-123 1-123 1-123 1-123 1-123 1-123 1-123	1 - 123 1 -
163 - 173 173 - 173 173 -	163 - 173 163 -	1
139 - 249 139 - 249 139 - 249 139 - 249 139 - 249 139 - 249 139 - 249	139 - 249 139 - 249	139 - 249 139 - 249
570 571 573 573 574 576 576	578 580 581 582 583 584 585	588 589 590 591 594 595 596 600 601 603
1090G09 1090G10 1090G12 1091A02 1091A03 1091A11	1091B02 1091B04 1091B05 1091B07 1091B11 1091B12 1091C02 1091C03	1091C05 1091C05 1091C06 1091C01 1091D01 1091D05 1091D05 1091E01 1091E01 1091E04 1091E04

	•		•		_
	SKULLLFPHDPLGF (SEQ ID NO SRDLLLFPHYGMDV (SEQ ID NO SRDLLLFPQSPLLF (SEQ ID NO: SRDLLLFPHEHLSF (SEQ ID NO: SRDLLLFPHSPLDF (SEQ ID NO:	SRDLLLFPHSPLSP (SEQ ID NO: 2 SRDLLLFPHYGMDV (SEQ ID NO SRDLLLFPNAALYP (SEQ ID NO: SRDLLLFPNDPLFG (SEQ ID NO: SRDLLLFPGAPLSP (SEQ ID NO: 2	ARDILLFPAAPLWP (SEQ ID NC SRDLLLFPNDPLR (SEQ ID NO: SRDLLLFPTAPLDP (SEQ ID NO: SRDLLLFPTAPLFP (SEQ ID NO: SRDLLLFPSDPLVF (SEQ ID NO: SRDLLLFPSDPLVF (SEQ ID NO:	SRDLLLFPGSPLTF (SEQ ID NO: SRDLLLFPYSHLEF (SEQ ID NO: SRDLLLFPQSPLHP (SEQ ID NO: SRDLLLFPQAPLFP (SEQ ID NO: SRDLLLFPQAPLFP (SEQ ID NO: SRDLLLFPYAPLTF (SEQ ID NO: SRDLLTF (SEQ ID NO: SR	SRDLLLFPQNPLHP (SEQ ID NO: 2506) SRDLLFPREPLCF (SEQ ID NO: 2636) SRDLLFPSAPLSF (SEQ ID NO: 2611) SRDLLFPMAPLRF (SEQ ID NO: 2593) SRDLLFPRSPLSF (SEQ ID NO: 2557) SRDLLFPRAPLYP (SEQ ID NO: 2387) SRDLLFPRDPLQF (SEQ ID NO: 2387) SRDLLFPTAPLTF (SEQ ID NO: 2531) SRDLLFPYSPLYP (SEQ ID NO: 2710) SRDLLFPYSPLYP (SEQ ID NO: 2710)
		1 1 1 1 1	1 1 1 1 1	1 1 1 1	99-112 99-112 99-112 99-112 99-112 99-112 99-112
		50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66		50 - 66 50 - 66
26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35	26-35 26-35 26-35 26-35 26-35		26 - 35 26 - 35
1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1-123	1-123 1-123 1-123 1-123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123
228 - 238 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238	1 1 1 1 1				228 - 28 228 - 28 228 - 28 228 - 28 228 - 28 228 - 28 228 - 28
189 - 195 189 - 195 189 - 195 189 - 195 189 - 195		1 4 1 1 1			189 - 195 189 - 195 189 - 195 189 - 195 189 - 195 189 - 195 189 - 195
163 - 173 163 -	21- 21- 21- 21- 21-	21- 21- 21- 21- 21- 21-	1.22 1.22 1.23 1.25 1.25 1.25 1.25 1.25 1.25 1.25 1.25	- 173 - 173 - 173 - 175 - 175 - 175	163 - 173 1 163 -
139 - 249 16 139 - 249 16 139 - 249 16 139 - 249 16 139 - 249 16		2242	2		249 16 249 16 249 16 249 16 249 16 249 16
139	139	139 139 139	139		139 - 9 - 9 - 9 - 9 - 9 - 9 - 9 - 9 - 9 -
606 607 609 609 611	613 614 615 616	618 620 620 621	625 625 626 626 627	628 629 630 631	633 637 637 638 639 640 640
1091E08 1091E09 1091E10 1091F01 1091F03 1091F05	1091F07 1091F08 1091F09 1091F10	1091F11 1091G01 1091G03 1091G04	1091G05 1091G06 1091G07 1091G09	1091G11 1091G12 1104A01 1104A07	1104A10 1104A10 1104A11 1104B02 1104B04 1104B11 1104C01

	SRDLLLFPRHPLLF (SEQ ID NO: 2543)							SRDLLLFPHYPLFI (SEO ID NO:	SRDLLLFPOAPLHP (SEC) ID NO	SRDLLLFPHAPMDP (SEO ID NO	SRDLLLFPRAPLTF (SEO ID NO:	SRDLLLFPRATLEF (SEO ID NO:	SRDLLLFPHSPLFP (SEO ID NO:	SRDLLLFPNDPLVL (SEQ ID NO	Ö	ON CI	SRDLLLFPASPLNP (SEO ID NO:	ON O	D NO	D NO:	SRDLLLFPHGPLTF (SEO ID NO:	SRDLLLFPHAPLSP (SEO ID NO: 2573)	SRDLLLFPSSPLIL (SEQ ID N		SRDLLLFPQDPLVF (SEQ ID NO: 2708)	SRDLLLFPKAPLVF (SEQ ID NO: 2544)			SRDLLLFPTAPLNF	SRHLLLFPQGPLSF	SRDLLLFPHLPLNP	SRDLLLFPHHSFDL (SEQ ID NO: 2147)	SRDLLLFPGAPLAP (SEQ ID NO: 2487)	SRDLLLFPQAPLYP (SEQ ID NO: 2378)	SRDLLLFPRSPLSF (SEQ ID NO: 2557)
	99 - 112	,	t				99 - 112	99 - 112	_		99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	7	-	•	7	99 - 112		99 - 112	99 - 112	$\overline{}$	99 - 112	99 - 112
	50 - 66	. ,		1		•	50 - 66	50 - 66		99 - 09	50 - 66	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	50 - 66	20 - 66	50 - 66	99 - 0 9	20 - 66	20 - 66	20 - 66	20 - 66				20 - 66	• •		•				20 - 66
	26 - 35		. •		- 1		26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35					<u>.</u>	٠,		26 - 35	26 - 35	26-35	26 - 35
	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123
	89 - 195 228 - 238 88 - 194 227 - 237	- 195 228-	-195 228-		189-195 228-238	189-195 228-238	189 - 195 228 - 238	189-195 228-238	189-195 228-238	189-195 228-238	-195 228-	189 - 195 228 - 238	-195 228-	- 195 . 228 -	1	-195 228-	189 - 195 228 - 238	- 195 228 -	- 195 228 -	- 195 228 -	-195 228-		-195 228-	-195 228-	- 195 228 -	- 195 228 -	-195 228-	- 195 228 -	- 195 228 -	- 195 228 -	- 195 228 -	- 195 228 -	- 195 228-	-195 228-	89 - 195 228 - 238
	163 - 173 1 162 - 172 1	- 173	- 173	-173. 1	163-173 1	163 - 173 1	- 173	-173 1	-173	- 173	-173	- 173	- 173	- 173	- 173	-173	- 173	- 173	-173	- 173	- 173	- 173	- 173	- 173	- 173	- 173	- 173	- 173	- 173	- 173	-173 1	-173	- 173	- 173	163 - 173 18
	139 - 249 138 - 248	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	•	•	139 - 249	•	139 - 249	139 - 249	139 - 249
;	6 42	44	645	. 949	647	648	£	· 620	651	652	653	654	655	959	657	658	629	099	199	995	993	664	999	999	299	899	699	0/9	671	672	673	674	675	9/9	677
	1104C05 1104C06	1104C07	1104C09	1104C11	·1104D01	1104D02	I104D03	I104D04	1104D07	1104D08	1104D09	1104E01	1104E02	1104E03	1104E05	1104E11	1104E12	1104F02	I104F03	1104F04	1104F05	1104F06	1104F07	1104F10	1104F11	1104F12	1104G04	1104605	1104G09	1104G11	1105A02	1105A03	1105A04	1105A08	1105A09

•			· ·	
SRDLLLFPSHSFDI (SEQ ID NO: 2692) SRDLLLFPYSPLHP (SEQ ID NO: 2658) SRDLLLFPYSPLSF (SEQ ID NO: 2676) SRDLLFPHHSFDL (SEQ ID NO: 2147) SRDLLFPHHSFDL (SEQ ID NO: 2147) SRDLLFPASPLNP (SEQ ID NO: 2364) SRDLLFPHEPLSP (SEQ ID NO: 2651) SRDLLFPHEPLSP (SEQ ID NO: 2651)		SRDLLLFPRDPLSF (SEQ ID NO: 2368) SRDLLLFPYAPLAF (SEQ ID NO: 2608) SRDLLLFPHAAFDV (SEQ ID NO: 2619) SRDLLFPHEPLFP (SEQ ID NO: 2640) SRDLLLFPHSALTF (SEQ ID NO: 2519) SRDLLLFPHHSFDS (SEQ ID NO: 2519) SRDLLLFPHHSFDS (SEQ ID NO: 2133)		SRDLLLFPKHPLVF (SEQ ID NO: SRDLLLFPHHSFDA (SEQ ID NO: SRDLLLFPHHSFDA (SEQ ID NO: SRDLLLFPHPPLLF (SEQ ID NO: SRDLLLFPKHPLVF (SEQ ID NO: SRDLLLFPKHPLVF (SEQ ID NO: SRDLLLFPKAPLYP (SEQ ID NO: SRDLLLFPKAPLYP (SEQ ID NO: SRDLLLFPHAPLDP (SEQ ID NO: SRDLLLFPHAPLP (SEQ ID NO: SRDLLLFPHAPLP (SEQ ID NO: SRDLLLFPHAPLP (SEQ ID NO: SRDLLTFPHAPLP (SEQ ID NO: SRDLTFPHAPLP (SEQ ID NO: SRDLTFPHAPL
99 - 112 99 - 112 99 - 112 99 - 112 99 - 112 99 - 112		99 - 112 99 - 112 99 - 112 99 - 112 99 - 112 99 - 112		
50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66		50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66		
26 - 35 26 - 35		26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35		
1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	1 - 123 1 - 123 1 - 123 1 - 123 1 - 123 1 - 123	
228 - 238 228 - 238		228 - 238 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238 228 - 238		
89 - 195 89 - 195 89 - 195 89 - 195 89 - 195 89 - 195	- 195 - 195 - 195 - 195 - 195 - 195	89 - 195 89 - 195 89 - 195 89 - 195 89 - 195 89 - 195		
163 - 173 1 163 - 173 1	173 - 173 -	21. 21. 21. 21. 21. 21. 21. 21. 21. 21.	21. 21. 21. 21. 21. 21. 21. 21. 21. 21.	
139 - 249 10 139 - 249 10	-249 -249 -249 -249 -249 -249	249 - 249 - 249 - 249 - 249		- 249 - 249 - 249 - 249 - 249 - 249 - 249
*				707 708 709 710 712 713
		7		
1105A11 1105B04 1105B05 1105B07 1105B10 1105B11 1105B12	1105C02 1105C03 1105C05 1105C06 1105C12 1105C12	1105D06 1105D08 1105D09 1105D10 1105D11 1105E01	1105E11 1105E11 1105F03 1105F06 1105F07 1105F12	1105G08 1105G09 1105G10 1105G11 1107A01 1107A03

										•																						
(SEQ ID NO:	SKULLIFFHAFLSF (SEQ ID NO: 2364) SRDLLIFPHAFLFP (SEQ ID NO: 2533)	SRDLLLFPASPLTF (SEQ ID NO: 2420)	SKULLLEFFHYGMUV (SEQ ID NO: 2133)		SRDLLLFPHYPLHP (SEQ ID NO: 2357)	_	(SEQ ID NO:	(SEQ ID NO:		(SEQ ID NO:	(SEQ ID NO:		(SEQ ID NO			SRDLLLFPKAPLTF (SEQ ID NO: 2382)	SRDLLLFPSAPLSP (SEQ ID NO: 2623)	(SEQ ID NO	(SEQ ID NO:	SRDLLLFPRTPLLF (SEQ ID NO:	SRDLLLFPNAPLSP (SEQ ID NO:	SRDLLLFPSAPLYP (SEQ ID NO:	SRDLLLFPHHSFDL	-	SRDLLLFPHYPLEM (SEQ ID	SRDLLLFPHAPLAP (SEQ ID NO:	SRDLLLFPHHSFDL (SEQ ID NO:	SRDLLLFPRDPLLF (SEQ ID NO:	SRDLLLFPLSPLVP (SEQ ID NO:	SRDLLLFPHDPLGF (SEQ ID NO:	SRDLLLFPHHSLLF (SEQ ID NO:	SRDLLLFPASPLNP (SEQ ID NO: 2364)
1 1	99 - 112 99 - 112	7	211 - 66	7	99 - 112		•			•	1			99-112		99 - 112		99 - 112	•	99 - 112	$\overline{}$	$\overline{}$	-	7	99 - 112	7	7		7	7	7	99 - 112
1 1	20 - 66 50 - 66	•	20-06			99-05								•	99 - 09	20 - 66		99 - 09	99 - 09	20 - 66	t		•			20 - 66				t	•	20 - 66
26 - 35	26 - 35 26 - 35					26 - 35	•	1			•	26 - 35		26-35	Ü	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35	٠,	26 - 35		26 - 35	26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35
1-123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123
1 1	228 - 238	•	228 - 238			228 - 238					•	228 - 238	•	228 - 238	228 - 238	228 - 238	228 - 238	228 - 238	228 - 238	•	228 - 238		228 - 238		228 - 238	228 - 238	228 - 238	t		•	228 - 238	228 - 238
1 1	189 - 195 189 - 195	· •	189 - 195		1	1	•	·			•	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	189 - 195	$\overline{}$	189 - 195	189 - 195	189 - 195	\mathbf{r}	189 - 195	7	189 - 195	189 - 195
77	163 - 173 163 - 173	$\overline{}$	163 - 173	7	7	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	[63 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173
139 - 249	139 - 249 139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	ŧ	139 - 249
714	716	718	719	721	727	723	724	725	726	727	728	729	730	731	732	733	734	735	736	737	738	739	740	741	742	743	74	745	746	747	748	749
1107A07 1107A09	1107A12 1107B02	1107B04	1107B05	1107001	1107C04	1107C06	1107C08	1107C10	1107D01	1107D04	1107D07	1107011	1107E01	1107E05	1107E07	1107E09	1107F01	1107F05	1107F09	1107F10	1107G01	1107G05	1107H02	· 1107H06	1107H09	1107H10	1108A12	1108B03	I108B04	1108C09	1108C11	1108D10

SPLNP (SEQ ID NO: 2364) API.NP (SEQ ID NO: 2709)	(SEQ ID NO:	(SEQ ID NO:	(SEQ ID NO:	APLDF (SEQ ID NO: 2369) VPI I F (SEQ ID NO: 2344)	(SEO ID NO:	(SEO ID NO	(SEQ ID NO:	_	(SEQ ID NO	_	APLAP (SEQ ID NO: 2381)	APLAP (SEQ ID NO: 2476)			_				SPLNP (SEQ ID NO: 2364)	APLHP (SEQ ID NO: 2691)		HGLDL (SEQ ID NO: 2449)	DPLLF (SEQ ID NO: 2515)	HSFDL (SEQ ID NO: 2147)		_	HSFDL (SEQ ID NO: 2150)	_		HGFDA (SEQ ID NO: 2703	APLWP (SEQ ID NO: 2352)	EPLAP (SEQ ID NO: 2434)	HPLEP (SEQ ID NO: 2411)
SRDLLLFPASPLNP SRDLLFPSAPLNP	SRDLLLFPHHSFDL	SRDLLLFPKHPLRF	SRDLLLFPHAPLFP	SRULLEFPHAFLUF	SRDLLLFPSAPLSP	SRDLLLFPRDPLDL	SRDLLLFPRDPLEF	SRDLLLFPNAPLSP	SRDLLLFPHYPFDA	SRDLLLFPRDPLRF	SRDLLLFPDAPLAP	SRDLLLFPRAPLAP	SRDLLLFPHHSLLF	SRDLLLFPHHPLTF	SRDLLLFPHHPLTF	SRDLLLFPRDPLHF	SRDLLLFPNAPLNP	SRDLLLFPHHSFDL	SRDLLLFPASPLNP	SRDLLLFPQAPLHP	SRDLLLFPHHSFDL	SRDLLLFPQHGLDL	SRDLLLFPRDPLLF	SRDLLLFPHHSFDL	SRDLLLFPQAPLHP	SRDLLLFPHYPLLF	SRYLLLFPHHSFDL	SRDLLLFPRAPLYP	SRDLLLFPKAPLDF	SRYLLLFPQHGFDA	SRDLLLFPSAP	SRDLLLFPQEPLAP	SRDLLLFPHHPLEP
99 - 112			99-112			7	99-112	99 - 112	99-112	.99 - 112		99-112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99 - 112	99-112	99 - 112	$\overline{}$	7	_	99 - 112	7	99 - 112	99-112	99-112	99-112	99 - 112	99 - 112	99 - 112	99 - 112
50 - 66	20 - 66	•	99 - 05	90 - 00 20 - 66	50 - 66	20 - 66	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	20 - 66	•		20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66
26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35
1 - 123 1 - 123	1-123	1 - 123	1-123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1-123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1-123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1 - 123	1-123
228 - 238	1		228 - 238			- 1	228 - 238	228 - 238	228 - 238	228 - 238		228 - 238	228 - 238	228 - 238		228 - 238	228 - 238		•	228 - 238		``	228 - 238	228 - 238		229 - 239	228 - 238	228 - 238	228 - 238	228 - 238	228 - 238	228 - 238	228 - 238
189 - 195 189 - 195	7	7	189 - 195	7	· ¬	7	7	189 - 195	189 - 195	⋾	7	189 - 195	7	189 - 195	$\overline{}$	189 - 195	189 - 195	-	∹	7	7	7	-	189 - 195	╗	189 - 195	189 - 195	⁻	7	7	7	189 - 195	189 - 195
163 - 173 163 - 173	7	163 - 173	163 - 173	7	7	7	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	$\overline{}$	7	163 - 173	163 - 173	7	7	•	•	$\overline{}$	₹	163 - 173	7	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173	163 - 173
139 - 249 139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249		139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249	139 - 249		•		139 - 250	•				139 - 249	139 - 249	139 - 249
750 751	752	753	754	756	757	758	759	760	761	762	763	764	765	992	167	298	769	770	771	2772	773	774	775	776	777	778	. 622	780	. 781	782	783	784	785
1108D11 1108D12	1108E01	1108E03	1108E05	1108E08	1108E09	1108E10	I108E11	1108F10	I108F12	I108G01	1108G02	1108G07	I108G10	1108G11	1108G12	1108H01	I108H02	1108H06	1108H08	· I111A06	1111B12	1111001	1111D06	1111E04	1111E10	1111E11	1111E12	1111F07	1111G02	1111H10	1113A04	I113A12	1113B06

54) 54) (0: 2172) 2171) 2171) 2171) 55) 2162) 2162) 2162)
8) 7) 7) 7) 4) 7) (4) (9) (9) (10
RDLLLFPHRFDL (SEQ ID NO: 24) RDLLLFPHRFDL (SEQ ID NO: 214) RDLLLFPHRFDL (SEQ ID NO: 215) RDLLLFPHRFDL (SEQ ID NO: 215) RDLLLFPHRFDL (SEQ ID NO: 215) RDLLLFPHRFDL (SEQ ID NO: 216) RRYDLLTFPHRFDL (SEQ ID NO: 216) RRYDLLTFPHRFDL (SEQ ID NO: 216) RRYDLLTGYYDNYWYFDL (SEQ ID NO: 216) RRYDLLTGYYDNYWYFDL (SEQ ID NO: 216) RRYDLLTGYYDNYWYFDL (SEQ ID NO: 216) RRYDLLTGYYDRYGWGWGW (SEQ ID NO: 216) RAYDPLTGYYRGHYFDY (SEQ ID NO: 216) RAYDPLTGYYRGHYFDY (SEQ ID NO: 216) RAYDPLTGYYFDGFDI (SEQ ID NO: 216) RAYDPLTGYYFGYFDI (SEQ ID NO: 216) RAYDPLTGYYFGYFDY (SEQ ID NO: 216) RAYDPLTGYYFDY (SEQ ID NO: 216) RAYDPLTGYYFDY (SEQ ID NO: 216) RAYDPLTGYYFDY (SEQ ID NO: 216)
SRDLLEPHHRFDL (SEQ ID SRDLLEPHHRFDL (SEQ ID SRDLLLEPHHSFDL (SEQ ID SRDLLLEPHYPLLF (SEQ ID SRDLLLEPHYPLLF (SEQ ID DRSYDILTGYYWYFDL (SEQ ID DRSYDILTGYYWYFDL (SEQ ID DRSYDILTGYYWYFDL (SEQ ID DRYDLLTGYYTDNYMDV (SEQ ID DRYDLLTGYYTDNYMDV (SEQ ID SRDLLLEPHYPDY GYYTDNYMDV (SEQ ID CRYDLLTGYYTGHLGYYTTDN (SEQ ID NG TYDPLTGYSFDGFDI (SEQ
SRDLLLFPHHRFDL SRDLLLFPHHSFDL SRDLLLFPHHSFDL SRDLLLFPHHSFDL SRDLLLFPHHRFDL SRDLLLFPHHRFDL SRDLLLFPHHRFDL SRDLLLFPHHSFDL SRDLLLFPHHSFDL SRDLLLFPHYPLLF SRDLLLFPHYPLLF SRDLLLFPHYPLTF SRDLLLFPHYPLTF SRDLLLFPHYPLTF SRDLLLFPHYPLTF SRDLLLFPHYPLTF SRDLLLFPHYPLTF SRDLLLFPHYPLTF SRDLLTFYYDL DGSYDLTGYYDD DRSYDLTGYYDD SHYDDLTGYYGG FNYDDLTGYSFDGF GRUDHYGNDL GYHDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF ATYDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF ATYDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF ATYDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF GGGNYDLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF GYHDPLTGYSFDGF GGGNYDLTGYSFDGF ATYDPLTGYSFDGF GGGNYDLTGYSFDGF ATYDPLTGYSFDGF
99 - 112 99 - 112 99 - 112 99 - 112 99 - 112 99 - 112 99 - 114 99 - 116 99 - 117 99 - 117 99 - 117
50 - 66 50
\$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$
1. 123 1.
228 - 238 - 248 -
89 - 195 89 - 195 80 - 195 80 - 195 80 - 196 80 - 1
173 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
163 - 163 - 164 - 165 -
249 249 249 249 249 249 249 249 249 249
139 - 139 - 139 - 139 - 139 - 139 - 139 - 139 - 139 - 139 - 139 - 141 -
823 824 825 826 827 827 831 831 833 833 834 840 841 844 844 845 845 855 855 853
1115H09 1116A07 1116B01 1116B01 1116E02 1116E02 1116E02 1116E02 1116F02 1100C09 1007H08 1007H08 1008B01 1008C02 1008C02 1008C02 1008C02 1008C02 1008C03

	<u>@</u>	6	<u>ر</u>																											
GGDYDILTGLYYYGMDV (SEQ ID NO: 2156) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) DGSYDILTGYYTDNYMDV (SEQ ID NO: 2154)	GYDSSAFKAFDI (SEQ ID NO: 2136) SSPPRWYDALTGDSSYHSAMDV (SEQ ID NO: 2169) DEGRDI I TGYYWPNEFDS (SEO ID NO: 2169)	SSPPKWYDALTGHSSYHSAMDV (SEQ ID NO: 2159)	SSPPKWYDALTGDSSYHSAMDV (SEQ ID NO: 2165)	DEGRDLLTGYYWPNFFDS (SEQ ID NO: 2168)	DGDILLVPAALMDV (SEO ID NO: 2160)		•	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)		ERHYYDILTGYQTGYGMDV (SEQ ID NO: 2784)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	BNO	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	DRETKVGYGMDV (SEQ ID NO: 2945)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	EGGNYDILTGYYIGNGAFDI (SEQ ID NO: 2158)	EGSYDIL TGYYVGVGRMDV (SEQ ID NO: 2171)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	DSYDILTGYRGYYFDY (SEQ ID NO: 2745)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	EGGNYDILTGYYIGNGAFDI (SEQ ID NO: 2158)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ERHYYDILTGYQTGYGMDV (SEQ ID NO: 2784)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ELGSSIVGATTGALDM (SEQ ID NO: 2852)	ELGSSIVGATTGALDM (SEQ ID NO: 2852)	DGYYDILTGYSYYGMDV (SEQ ID NO: 2135)	MEYDIL TGYYGGYFDY (SEQ ID NO: 2179)
99 - 115 99 - 114 99 - 116	99 - 120			99 - 116	99 - 113	104 - 118	102 - 117		101 - 119	99 - 117		99 - 114	99 - 114	99 - 114	99 - 110	99 - 114	97 - 116	99 - 117	99 - 114	98 - 113	99 - 114	97 - 116	99 - 114	99 - 114	99 - 117	99 - 114	98 - 113	98 - 113	99 - 115	99 - 114
50 - 66 50 - 66 50 - 66	. 50 - 66 50 - 66	50 - 66	20 - 66	50 - 66	20 - 66	55 - 71	52 - 69	20 - 66	20 - 68	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	48 - 64	20 - 66	50 - 66	50 - 65	20 - 66	48 - 64	50 - 66	99 - 09	99 - 05	99-09	49 - 65	49 - 65	20 - 66	20 - 66
26 - 35 26 - 35 26 - 35	26 - 35	26 - 35	26 - 35	26-35	26-35	26 - 40	26-37	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	24 - 33	26 - 35	26 - 35	26 - 35	26 - 35	24 - 33	26 - 35	26-35	26 - 35	26-35	26 - 34	26 - 34	26 - 35	26-35
1-126 1-125 1-127	1-121	1-131	1-131	1-127	1 - 124	1 - 129	1 - 128	1 - 125	1 - 130	1 - 128	1 - 125	1-125	1 - 125	1 - 125	1 - 121	1 - 125	1 - 127	1 - 128	1-125	1 - 124	1 - 125	1 - 127	1 - 125	1 - 125	1 - 128	1 - 125	1 - 124	1 - 124	1 - 126	1 - 125
231 - 242 231 - 239 232 - 242	238 - 247 238 - 247 234 - 244	237 - 248	237 - 247	234 - 244 230 - 239		236 - 246	234 - 243	230 - 240	236 - 245	234 - 243	•	231 - 240	231 - 240	227 - 237	226 - 236	231 - 240	233 - 242	234 - 243	231 - 240	•	231 - 240	233 - 242	231 - 240	231 - 240	234 - 243	,	1			227 - 237
	- 205	-204	- 204	- 201		-203 2	-201			-201 2	- 198		192 - 198 2	- 194	- 193	- 198	200	201	- 198			- 200	• •	- 198 2	201	• •	•		• •	- 194 2
192	199	198	198	195	161	197	195	191	197	195	192	192	192	188	187	192	194-	195-	192	191	192	194	192	192	195	192	192	192	189	188
164 - 176 163 - 176 165 - 177	100 - 172 170 - 183 167 - 179	•	70 - 182	167 - 179 162 - 175	162 - 175	68 - 181	166 - 179	$\overline{}$	7	166 - 179	1	7	163 - 176	$\overline{}$	7	163 - 176	•	$\overline{\cdot}$	7	62 - 175	163 - 176	165 - 178	163 - 176	163 - 176	166 - 179	163 - 176	163 - 176	163 - 176	63 - 173	62 - 172
253 250 253	- , ,	_	-258 1	-255	250	-257 1	-254	-251	-256	- 254	_	Ξ.	-251 1	- 248 1	-247 1	- 251	- 253	- 254	-251	- 250	-251 1	-253 1	-251 1	-251 1	-254 1	· .	-251	-251	- 249 1	-248
142- 141: 143-	147	147	147	143	140	145	144	141	146	144	141	141	141	141	137	141	143	4	141	140	141	143	141	141	144	141	140	140	142	141
858 859 860	862 863	864	865	866 867	898	698	870	871	872	873	874	875	876	877	878	879	. 880	881	882	883	884	885	886	887	888	886	880	891	892	893
1028A06 1029D07 1029F11	1031C07 1031F09	I031G08	1031G10	1031G11 1037E07	1037E12	I050A07	I061D02	I061E07	I061HI01	I001A03	I001A07	I001A08	I001A10	I001A12	I001B02	I001B07	1001C06	I001C08	I001C12	1001D08	I001D12	I001E05	I001E07	I001G09	I001H05	I001H08	I003A01	I003A06	I003A07	I003A10

	~	•	
ELGSSIVGATTGALDM (SEQ ID NO: 2852) RYGDPFYYYYMNV (SEQ ID NO: 2755) DGYYDILTGYSYYGMDV (SEQ ID NO: 2135) ELGLSIVGATTGALDM (SEQ ID NO: 2174) GDYDILTGYPAECFQI (SEQ ID NO: 2854) GDYDILTGYPAECFQI (SEQ ID NO: 2854) MEYDILTGYYGGYFDY (SEQ ID NO: 2179) RYGDPFYYYYYMNV (SEQ ID NO: 2755) GDYDILTGYPAECFQI (SEQ ID NO: 2755)		SHYDILTGLNYWYFDL (SEQ ID NO: 2166) EGRDILTGVYYYGLDV (SEQ ID NO: 2893) SHYDILTGLNYWYFDL (SEQ ID NO: 2166) TYYDILTGRFFDI (SEQ ID NO: 2166) SHYDILTGRFFDI (SEQ ID NO: 2166) DLRYDILTGYHDAFDI (SEQ ID NO: 2166) DLRYDILTGYYFYGMDV (SEQ ID NO: 2890) GTYYDILTGYYFYGMDV (SEQ ID NO: 2774) SHYDILTGLNYWYFDL (SEQ ID NO: 2166) SHYDILTGLNYWYFDL (SEQ ID NO: 2166) DQHDILTGVYYGMDV (SEQ ID NO: 2166)	VSPSYDILTGYYLPHAFDV (SEQ ID NO: 2849) TYYDILTGRFFDI (SEQ ID NO: 2866) PSYDILTGYLYYFDY (SEQ ID NO: 2850) DLRYDILTGYYHDAFDI (SEQ ID NO: 2850) GAYYDILTGYYPYGMDV (SEQ ID NO: 2860) GQYYDILTGYYWYPYGMDV (SEQ ID NO: 2857) SRDILLFPHYGMDV (SEQ ID NO: 2857) SRDILLFPHYGMDV (SEQ ID NO: 2133) GGYSSGWLRGGPYNWFDP (SEQ ID NO: 2957)
98 - 113 99 - 115 99 - 114 99 - 114 99 - 114 99 - 114		99 - 114 99 - 114 99 - 114 99 - 114 99 - 116 99 - 116 99 - 117 99 - 115 99 - 115 99 - 115 99 - 115 99 - 115 99 - 113	99 - 117 98 - 110 99 - 113 99 - 115 99 - 115 99 - 112
49 - 65 50 - 66 49 - 65 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 65 50 - 65 50 - 66 50 - 66 50 - 66 50 - 66
26 - 34 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35
1-124 1-126 1-126 1-127 1-125 1-125 1-125 1-125	1-124 1-123 1-127 1-129 1-128 1-124 1-124	1 - 125 1 - 125 1 - 125 1 - 121 1 - 125 1 - 126 1 - 126 1 - 126	1 - 128 1 - 121 1 - 124 1 - 125 1 - 126 1 - 124 1 - 123
231 - 240 227 - 237 228 - 238 231 - 241 231 - 241 229 - 239 227 - 237 229 - 239		227 - 238 227 - 237 227 - 237 227 - 236 227 - 237 231 - 240 230 - 238 230 - 238 227 - 237	232 - 240 227 - 236 226 - 236 231 - 240 230 - 238 226 - 236 225 - 235 232 - 242
198 198 196 196 196	201 201 203 203 203 198 198	••••	199 194 193 197 193 199
176 192- 173 188- 173 189- 176 192- 174 190- 174 190- 176 192- 176 192-		44444668864	
_ , , , , , , , , , ,	163 - 1 163 - 1 162 - 1 168 - 1 168 - 1 164 - 1 163 - 1	162 - 17 162 - 17 162 - 17 159 - 17 162 - 17 163 - 17 165 - 17 165 - 17	167-1 159-1 161-1 163-1 165-1 160-1
- 251 - 248 - 249 - 252 - 250 - 250 - 250 - 250		- 248 - 248 - 248 - 248 - 249 - 249 - 249	
140 138 142 140 141 141 141 139	139 139 143 143 144 140 140	141 141 137 141 142 142 144 140	144 137 140 141 142 140 139
894 895 896 897 898 899 900 901	903 904 905 905 906 910	912 913 914 915 917 918 920 920	22 22 22 22 22 22 22 22 22 22 22 22 22
1003B03 1003B04 1003B09 1003C01 1003C02 1003C12 1003D04 1003B05	1003F01 1003F01 1003G01 1003G05 1003G06 1003H02 1003H02	1005A01 1005A02 1005B01 1005B09 1005C01 1005D02 1005E01 1005E01 1005E01	1005F02 1005F04 1005F08 1005G01 1005G08 1005H02 1006B01

				•	
GDYDILTGYYPLRDY (SEQ ID NO: 2792) NLFDVWTLPYYYYMDV (SEQ ID NO: 2965) ADYDILTGYSPLTYGMDV (SEQ ID NO: 2762) MYYDILTGHNFDY (SEQ ID NO: 2879) VSRDILTGHNFDY (SEQ ID NO: 2817) GGYSSGWLRGGPYNWFDP (SEQ ID NO: 2967) AGGYYDILTGRDYYYGMDV (SEQ ID NO: 2967) RRYALDY (SEQ ID NO: 2920)	DRGSYDILTGYYTPPHYYGMDV (SEQ ID NO: 2761) GGYSSGWLRGGPYNWFDP (SEQ ID NO: 2967) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) SHYDILTGLNYWYFDY (SEQ ID NO: 2746) ENYDFLTGYYGAFDI (SEQ ID NO: 2772) GRYDILTGYHWDGAFDI (SEQ ID NO: 2892)	ATYDFLIGYSFUGFDI (SEQ ID NO: 2133) ATYDFLTGYSFDGFDI (SEQ ID NO: 2153) IRLYCYSLTGYYPYGMDD (SEQ ID NO: 2810) TNYDILTGYYQGVDY (SEQ ID NO: 2782) GOYYDILTGYNWFDP (SEQ ID NO: 2857)	GRYDILTGYHWDDAFDI (SEQ ID NO: 2872) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) DFYDILTGYPLGGMDV (SEQ ID NO: 2741) DLPYYDILTGYSLTSGMDV (SEQ ID NO: 2923) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) GRRYDILTGYYYYHHGMDV (SEQ ID NO: 2811)	SHYDILTGLNYWYFDL (SEQ ID NO: 2166) DSGGDILTGYYMPYFDY (SEQ ID NO: 2847) VGLYYDILTGYYPSGMDV (SEQ ID NO: 2805) SQAHYDILTGYYLWSYGMDV (SEQ ID NO: 2875) ESYDILTGYRHYGMDL (SEQ ID NO: 2891) A TYNDILTGYSHYGMDL (SEQ ID NO: 2891)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) DREYDLLTGYYLHAFDM (SEQ ID NO: 2960) ENYDFLTGYYGAFDI (SEQ ID NO: 2772) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)
	98 - 119 99 - 116 99 - 114 99 - 113 101 - 117	99 - 114 99 - 115 99 - 113 99 - 113	101 - 117 99 - 114 99 - 117 99 - 117 99 - 116		99 - 114 99 - 114 99 - 115 99 - 115
50 - 66 50 - 68 50 - 68 50 - 68 50 - 66 50 - 66	50 - 65 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 65 50 - 65 50 - 66	50 - 68 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 68 50 - 68 50 - 68	
26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35 26-35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35		786-58
1 - 125 1 - 127 1 - 127 1 - 124 1 - 126 1 - 127 1 - 130	1 - 130 1 - 127 1 - 125 1 - 123 1 - 124 1 - 128	1 - 125 1 - 125 1 - 126 1 - 124	1 - 128 1 - 125 1 - 125 1 - 128 1 - 125	1-125 1-126 1-126 1-126 1-131	1 - 125 1 - 125 1 - 126 1 - 126 1 - 125
		231 - 240 231 - 240 228 - 238 230 - 239 226 - 236	230 - 240 227 - 237 231 - 240 230 - 240 227 - 237	231 - 240 232 - 241 228 - 238 237 - 246 227 - 237	
	193 - 199 193 - 199 192 - 198 190 - 196 191 - 197	192 - 198 192 - 198 189 - 195 191 - 197 187 - 193			
-176 -178 -176 -175 -177 -179	-177 -176 -176 -175	163 - 176 1 163 - 176 1 163 - 173 1 162 - 175 1	175	- 176 - 177 - 173 - 182 - 172	163 - 176 163 - 176 163 - 176 163 - 173 162 - 175
-251 -253 -250 -250 -253 -253 -253	-253 -253 -251 -249 -250	-251 -251 -249 -250	-251 -248 -251 -251 -248	252 252 252 2549 257	41 - 251
	938 146 939 143 940 141 941 139 942 140				960 14 962 17 963 14 964 16
1006D09 1006E01 1006E07 1006F01 1006F02 1006G01 1006G04	1006H01 1006H02 1007A01 1007A11 1007A11	1007B04 1007C04 1007C08 1007C12	1007E03 1007E03 1007E10 1007E11 1007F06	1007,503 1007,609 1007,610 1007,410	1008A02 1008A05 1008A06 1008A12 1008B02

							~																												
GSVDII TGVVIDNVAMDV (SEO ID NO: 2154)	DHYDILTGLYYYGMDV (SEO ID NO: 2760)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	GRRYDILTGYYKGPLDY (SEO ID NO: 2902)	AYYDNLTGFLPYGMGV (SEO ID NO; 2947)	EGYDIL TGYFLDYYHGMDV (SEO ID NO: 2753)	ATYDPLTGYSFDGFDI (SEO ID NO; 2153)	GPRGGPYYDILTGYYLSLSDAFDI (SEO ID NO: 2729)	EYYDIL TGYRDPYGMDV (SEO ID NO: 2973)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	EVRNYDLLTRSYLAGPLDN (SEO ID NO: 2751)		ATYDPLTGYSFDGFDI (SEQ ID NO; 2153)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	DRGYYDILTGYYRGHGMDV (SEO ID NO: 2837)	DLPYYDILTGYSLTSGMDV (SEQ ID NO: 2923)	EEGFYDILTGYYGPGYFDY (SEO ID NO: 2974)	ATYDPLTGYSFDGFDI (SEQ ID NO; 2153)	EGYDILTGYSKFLDY (SEQ ID NO: 2906)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	SHYDILTGLNYWYFDL (SEQ ID NO: 2166)	ERADYDIL TGYYFYDMDV (SEQ ID NO; 2833)	FRYDILTSYYYGMDV (SEQ ID NO: 2734)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	GRRYDILTGYYYYHHGMDV (SEQ ID NO: 2811)	GHYDIL TGYDDYYYGMDV (SEQ ID NO: 2844)	HDILTGFDY (SEQ ID NO: 2904)	SGYDILTGYLYGMDV (SEQ ID NO: 2934)	APYDILTGYSDYYGMDV (SEQ ID NO: 2968)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	GDYDPLTGYSFGVDV (SEQ ID NO: 2941)	EGSYDILTGYYVGVGRMDV (SEQ ID NO: 2171)	DGYYDILTGGFYYYYGMDV (SEQ ID NO: 2899)
116 1	115	114	113	114	117	114	22	115	14	114	117	114	114	114	117	-117	117	- 114	110	114	114	114	99-116 ERA	100 - 114 FRY	114	114	116	116	90	113	-1117	114	113	116	-117 DGY
00 99-	8		Z	- 66 99 - 05	-66 99-05	-66 99-05		50 - 66 99 - 1			50-66 99-1	-66 99-09	20-66 99-05	20-66 99-05	20-66 99-05	66 99-09	-66 99-09	66 99-05	46-63 96-	-66 99-05	-66 99-09	50-66 99-1	66 99-05						50-65 98-1	-66 99-05	50-68 101				.66 99-
08 - 36 - 90	-35		33	26-35 50	26-35 50	26-35 50	26-35 50	26-35 50	26-35 50	26-35 50	26-35 50	26-35 50	26-35 50	26-35 50	35	35	26-35 50	26-35 50	20-31 46	26-35 50	26-35 50	26-35 50	35	37		35	35	35	26-35 50	35	35	35	35	-35	26-35 50
1 - 127				1 - 125	1 - 128	1 - 125	1 - 133	1 - 126	1 - 125	1-125	1-128	1 - 125	1-125	1-125	1 - 128		1 - 128	1-125	1-121	1 - 125	1-125	1 - 125	1 - 127			1 - 125			1-117						1 - 128
233 - 242		- 1	228 - 236	227 - 237	234 - 243	231 - 240	238 - 248	228 - 238	231 - 240	231 - 240	234 - 243	231 - 240	231 - 240	227 - 237	234 - 243	230 - 240	234 - 243	227 - 237	227 - 236	231 - 240	227 - 237	230 - 241	233 - 242		•	231 - 240	•		223 - 232		230 - 240	231 - 240	226 - 236	•	234 - 243
194 - 200		192 - 198	189 - 195	188 - 194	195 - 201	192 - 198	199 - 205	189 - 195	192 - 198	192 - 198	195 - 201	192 - 198	192 - 198	188 - 194	195 - 201	191 - 197	195 - 201	188 - 194	188 - 194	192 - 198	188 - 194	191 - 197	194 - 200		192 - 198	192 - 198	1	194 - 200	184 - 190			•	187 - 193	•	195 - 201
165 - 178	162-	163 - 176	163 - 173	162 - 172	166 - 179	163 - 176	171 - 183	163 - 173	163 - 176	163 - 176	166 - 179	163 - 176	163 - 176	162 - 172	166 - 179	7	166 - 179	162 - 172	159 - 172	7	162 - 172	•	$\overline{}$	ï	7	163 - 176	7	165 - 178	155 - 168	161 - 171	165 - 175	163 - 176	161 - 171	165 - 178	166 - 179
143 - 253	•	141 - 251	140 - 247	141 - 248	144 - 254	141 - 251	149 - 259	142 - 249	141 - 251	141 - 251	144 - 254	141 - 251	141 - 251	141 - 248	144 - 254	144 - 251	144 - 254	141 - 248	137 - 247	141 - 251	141 - 248	141 - 252	143 - 253	•	•	•	•	•	133 - 243	•		•	140 - 247	143 - 253	144 - 254
996	296	896	696	970	126	972	. 973	974	975	926	716	978	979	086	981	982	983	984	985	986	282	886	686	986	166	992	993	994	995	966	997	866	666	1000	1001
I008B04	I008B05	I008B06	I008B07	I008B10	I008B11	1008C06	1008C08	I008C09	I008D01	I008D02	I008D03	I008D04	I008D05	. 90C800I	I008D07	I008D08	I008D12	I008E01	I008E02	I008E03	I008E04	I008E08	I008E09	I008E12	I008F03	I008F06	I008F07	I008F08	I008F09	I008F10	I008F11	I008G02	I008G03	I008G04	1008G05

			·	5
AYYDILTGLDY (SEQ ID NO: 2966) DQQYDILTGYYHYGMDV (SEQ ID NO: 2964) DQVDLLLMDHNYYMDV (SEQ ID NO: 2918) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) EGSYDILTGYYYVGVGRMDV (SEQ ID NO: 2171) DQQYDILTGYYYYYMDV (SEQ ID NO: 2964) TKYDILTGYYYYYMDV (SEQ ID NO: 2964) TKYDILTGYYYYYMDV (SEQ ID NO: 2174) ELGLSIVGATTGALDM (SEQ ID NO: 2174) ELGLSIVGATTGALDM (SEQ ID NO: 2174)	TDREGAKDVTSRWGMDV (SEQ ID NO: 2814) BLGLSIVGATTGALDM (SEQ ID NO: 2174) BLGLSIVGATTGALDM (SEQ ID NO: 2174) DRGGNYDILTGYYFHHGVDV (SEQ ID NO: 2914) BLGLSIVGATTGALDM (SEQ ID NO: 2174) RYGDPFYYYYYMNV (SEQ ID NO: 2755)	ELGLSIVGATTGALDM (SEQ ID NO: 2174) RYGDPFYYYYYMNV (SEQ ID NO: 2755) RYGDPFYYYYYMNV (SEQ ID NO: 2755) ELGLSIVGATTGALDM (SEQ ID NO: 2174) SSPPKWYDALTGHSSYHSAMDV (SEQ ID NO: 2159) SSPPKWYDALTGHSSYHSAMDV (SEQ ID NO: 2159)	ÖZÖZ ÖZ BABA B	GLRHVTLFGTGTRGHFYMDV (SEQ ID NO: 2789) GREDTDKVKPWDRYYHYYYMDV (SEQ ID NO: 2809) GYDSSAFRAFDI (SEQ ID NO: 2136) SSPPKWYDALTGDSSYHSAMDV (SEQ ID NO: 2165) GYDSSAFRAFDI (SEQ ID NO: 2136)
99 - 109 99 - 114 99 - 114 99 - 116 99 - 116 99 - 113 98 - 113	99 - 115 98 - 113 99 - 118 99 - 118		99 - 120 99 - 120 99 - 120 99 - 110 99 - 110	99 - 118 99 - 120 99 - 110 99 - 120 99 - 110
50 - 66 50 - 6	50 - 66 49 - 65 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66
26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 34 26 - 34 26 - 34 26 - 34 26 - 34 26 - 34 26 - 35	26-35 26-34 26-34 26-35 26-35	26 - 34 26 - 34 26 - 34 26 - 34 26 - 34 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 26 26 26 26 26 26 26 26 26 26 26 26	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35
1 - 120 1 - 127 1 - 125 1 - 127 1 - 127 1 - 127 1 - 124	1 - 126 1 - 124 1 - 124 1 - 129 1 - 123	1 - 124 1 - 124 1 - 123 1 - 123 1 - 131 1 - 131	1-131 1-131 1-131 1-121 1-121 1-121	1 - 129 1 - 131 1 - 121 1 - 131
226 - 235 233 - 242 229 - 237 231 - 240 233 - 242 233 - 242 229 - 237 230 - 238		228 - 238 228 - 238 231 - 240 228 - 239 230 - 240 230 - 240 228 - 238 237 - 248	237 - 245 236 - 246 237 - 247 228 - 238 228 - 237 238 - 248	236 - 246 238 - 248 228 - 238 238 - 249 227 - 237
7 - 193 60 - 196 7 - 198 7 - 198 7 - 198 9 - 196 1 - 197 1 - 197	4 - 200 2 - 198 0 - 196 6 - 202 2 - 198 0 - 196	198 198 197 197 197	8 - 204 7 - 203 8 - 204 9 - 195 9 - 205	197 - 203 199 - 205 189 - 195 199 - 205 188 - 194
171 178 174 176 178 178 178 178 175	- 178 1 - 176 1 - 174 1 - 180 1 - 176 1	173 176 177 173 173 173 182	- 182 - 182 - 173 - 173 - 183	-181 -183 -173 -183
		247 103 249 163 251 164 252 164 250 163 249 163 259 170 255 170		257 168 259 170 249 162 260 170 248 160
	142 - 2 140 - 2 140 - 2 140 - 2 130 - 2	140 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 -		145 - 2 147 - 2 137 - 2 147 - 2 137 - 2
1002 1003 1004 1005 1006 1007 1009 1010	1012 1013 1014 1015 1016	1020 1022 1022 1023 1023 1025	1027 1028 1029 1030 1031	1033 1034 1035 1036
1008G11 1008G12 1008H02 1008H03 1008H09 1008H11 1012B06	1012C03 1012C06 1012C09 1012D12 1012E07 1012E08	1012E03 1012F05 1012G03 1012G05 1012G10 1012G10	1013B04 1013B09 1013C02 1013C04 1013D02	1013D10 1013E02 1013E05 1013E09 1013F03

AMDV (SEO ID NO: 2159	VDV (SEQ ID NO: 2131)	40: 2136)	AMDV (SEQ ID NO: 2159	XXXIIIV (SEQ ID NO: 2159)	1 I'M (3EQ M NO. 28)		O ID NO: 2153)	O ID NO: 2153)	(C)) ID NO: 2757)	N (SEO ID NO: 2751)	2153)	O ID NO: 2153)	D NO:	YYFDY (SEO ID NO: 2132)		YYFDY (SEO ID NO: 2132	O ID NO: 2153)	D NO	ON CO	O ID NO: 2153)	D NO	D NO:	(SEQ ID NO: 2153)	(SEQ ID NO: 2153)	YYFDY (SEQ ID NO: 2132)	(SEQ ID NO: 2153)	O ID NO: 2153)	D NO:	(SEQ ID NO: 2157)	(SEQ ID NO: 2153)	B NO:	D NO:	ĊN
SSPPKWYDALTGHSSYHSAMDV (SEO ID NO: 2159)	AATTSQKHNKYAYYFYGMDV (SEQ ID NO: 2131)	GYDSSAFRAFDI (SEQ ID NO: 2136)	SSPPKWYDALTGHSSYHSAMDV (SEQ ID NO: 2159	GREDITOKVKPWDRYYHYVYYAMV (SEQ ID NO: 2139)	EGGNYDILTGYYIGNGAFDI (SEO ID NO:	GDYDILTGYPAECFOI (SEO ID NO: 2854)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	ATYDPL TGYSFDGFDI (SEO ID NO: 2153	ELGLSIVGATTGALDM (SEO ID NO: 2174	AGYDLLTGYPFYFDS (SEO ID NO: 2757)	EVRNYDLLTRSYLAGPLDN (SEO ID NO:	ATYDPLTGYSFDGFDI (SEO ID NO.	ATYDPLTGYSFDGFDI (SEO	ATYDPLTGYSFDGFDI (SEO ID NO: 2153	VQMDSEYYDLLTGINVGPYYFDY (SEO ID	ATYDPLTGYSFDGFDI (SEO	VOMDSEYYDLLTGINVGPYYFDY (SFO ID	ATYDPLTGYSFDGFDI (SEO	ATYDPLTGYSFDGFDI (SEO			ATYDPLTGYSFDGFDI (SEO	ATYDPLTGYSFDGFDI (SE	ATYDPLTGYSFDGFDI (SE	ATYDPLTGYSFDGFDI (SE(ATYDPLTGYSFDGFDI (SE(ATYDPLTGYSFDGFDI (SEQ)	ATYDPLTGYSFDGFDI (SEQ)	ATYDPLTGYSFDGLDI (SEC	ATYDPLTGYSFDGFDI (SEC	ATYDPLTGYSFDGFDI (SEQ)	ATYDPLTGYSFDGFDI (SEO	ATYDPLTGYSFDGFDI (SEO ID
99 - 120	99 - 118		99 - 120			99 - 114	99 - 114	99 - 114	98 - 113	100 - 114	99 - 117	99 - 114	99 - 114	99 - 114		99 - 114	t	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	99 - 114	•	99 - 114	99 - 121	99 - 114	99 - 114	99 - 114	98 - 113	99 - 114	99 - 114	99 - 114	99 - 114
50 - 66	20 - 66	50 - 66	20 - 66	20 - 66	48 - 64	20 - 66	20 - 66	50 - 66	49 - 65	52-67	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66		99 - 09	20 - 66	20-66	99 - 09	99 - 09	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	20 - 66	99-05		20 - 66	20 - 66	20 - 66	50 - 66
26 - 35			26 - 35			26 - 35	26-35			26 - 37	26 - 35	26 - 35	26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35		25 - 34	26 - 35	26 - 35	26-35	26 - 35
1-131	1 - 129	1 - 121	1 - 131		1-127	1 - 125	1 - 125	1 - 125	1 - 124	1 - 125	1 - 128	1 - 125	1 - 125	1 - 125	1 - 132	1 - 125	1 - 132	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 132	1 - 125	1 - 125	1 - 125	1 - 124	1 - 125	1 - 125	1 - 125	1 - 125
237 - 247	•		238 - 248	- (233 - 242	233 - 243	231 - 240	231 - 240	231 - 241	231 - 240	230 - 240	231 - 240	231 - 240	231 - 240	•	231 - 240	234 - 244	231 - 240	231 - 240	٠	•	231 - 240	•	•	1.		•		•	•		•	1 - 240	1-240
204 2			2042	205	200	200 2	198	198	198	198	197	198	198	198	201	198	201	198	198		٠.	861	861	86	861	50	86	861	861	192	86	198	198 231	198 231
198 -	201 -	188	198	199	194-	193-	192	192-	192.	192-	191	192	192-	192 -	195-	192 -	195-	192 - 198	192	192 - 198	192 - 198	192 - 198	192	192	192	195	192 - 1	192-	192-	191 -	192 -	192	192-	192 -
- 182	- 185	- 172	- 182	- 183	- 178	- 177	- 176	- 176	- 176	- 176	- 175	- 176	- 176	- 176	- 179	- 176	- 179	- 176	- 176	- 176	- 176	- 176	- 176	- 176	- 176	- 179	- 176	- 176	- 176	- 175	- 176	- 176	- 176	- 176
			2 2	170	165	164	163	163	164	166	165	163	163	163	169	163	169	163	163	163	163	163	163	163	163	169	163	163	163	162	163	163	163	163
•	•	137 - 248	. ,	•	143 - 253	141 - 254	141 - 251	141 - 251	140 - 252	141 - 251	144 - 251	141 - 251	•	1	٠	141 - 251	148 - 255	•	141 - 251	٠	•	1	•	1	•	t	•	٠	•	•	. 1	1	141 - 251	141 - 251
1038	1039	1040	1042	1043	1044	1045	1046	1047	1048	1049	1050	1051	1052	1053	1054	1055	1056	1057	1058	1059	1060	1061	1062	1063	1064	1065	1066	1067	1068	1069	1070	1071	1072	1073
1013F04	I013F07	1013F09 1013E10	1013H04	1013H07	I014A12	I014C06	I014C10	I014C12	I014E06	.1014F02	I016A08	I016A09	I016C02	I016C03	1016C05	1016C09	1016C11	I016D10	1016D111	1016E03	1016E04	I016F03	I016F11	1016G01	1016506	1016G12	1016H10	1017A06	I017A07	I017A11	1017E12	1017G03	1017G07	1017H01

						•																											
		ATVDN TOVEROFFI (SEQ	ATVIDITENSEDGEDI (SEQ ID NO:	ATVIDITION TO VED TO NO.	ATYDEI TGYSEDGEDI (SEQ ID NO:	ATVIDITEST STEPS (SECTION NO.	ATVIDE TRYSEDGEDI (SEQ ID NO:	ATVIDITIONED CENTROL (SECTION NO.	`.	ATYDPI TGYSEDGEDI (SEC ID NO:	ATYDPI.TGYSFDGFDI (SEO ID.	ERHYYDII.TGYOTGYC	ERHYYDILTGYOTGYGMDV (SEO ID NO:	EGGNYDII TGYYIGNGAEDI			EGGNYDII.TGYYTGNGAFDI (SEO ID NO.	EGGNYDILTGYYIGNGAFDI (SEO ID NO:	EGGNYHILTGYYIGNGAFDI (SEO ID NO:	EGENYDIL TGYYJGNGAFDI (SEO ID NO:	EGGNYDILTGYYIGNGAFDI (SEQ ID NO: 2158)	EGGNYDILTGYYIGNGAFDI (SEQ ID NO: 2158)	EGGNYDILTGYYIGNGAFDI (SEQ ID NO: 2158)	DRETKVGYGMDV (SEQ ID NO: 2945)	EGGN Y DILLIGY Y IGNGAFDI (SEQ ID NO: 2158)	EGGNYDILIGYYIGNGAFDI (SEQ ID NO: 2158)	DGYYDLTGYSYYGMDV (SEQ ID NO: 2135)	ATYDPLIGYSFDGFDI (SEQ ID NO: 2153)	MEYDIL TGYYGGYFDY (SEQ ID NO: 2179)	DGYYDILTGYSYYGMDV (SEQ ID NO: 2135)	AT Y DPL 1G Y SFDGFDI (SEQ ID NO. 2153)	ALTURIOR SECTION (SECTION) (2153)	ASYYDILTGYYKGAFDI (SEQ ID NO: 2135)
	99 - 117	• (99 - 114		-						97 - 116	97 - 116	97 - 116	97-116			27-110				•		99 - 114		99-115
	50 - 66 50 - 66		50 - 66	20 - 66	20 - 66	20 - 66	50 - 66	20 - 66	20 - 66	50 - 66	20 - 66	99 - 09	99-09	48 - 64	99-09	48 - 64	48 - 64	48 - 64	48 - 64	48 - 64	48 - 64	48 - 64	48-64	20 - 66	40-04	\$ - 05 5 - 05	00-00	20-00	20 - 66	90-00	90-00	00 - 00 80 - 66	99 - 05 20 - 66
	26 - 35	, ,	26 - 35	26 - 35	26 - 35	, (7)		26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	24 - 33	26 - 35	24 - 33	24 - 33	24 - 33	24 - 33	24 - 33		24 - 33	24 - 33	26 - 35	24 - 33	26 - 47			26 - 35	26 - 35	•	26-35	26 - 35
1 - 125	1 - 128	1-125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 128	1 - 128	1 - 127	1 - 121	1-127	1 - 127	1 - 127	1 - 127	1 - 127	1 - 127	1 - 127	1 - 127	1 - 121	1 - 127	121 - 1	1 - 120	751 - 1	1-125	1 - 120	1 - 125	1-126	1 - 126
•	234 - 243			- 1	•			231 : 240		231 - 240	231 - 240	234 - 243	234 - 243	233 - 242		233 - 242	233 - 242	•	•	t	t			220 - 230) (1		•	157 - 677				•
- 198	192 - 201	- 198	- 198	- 198	- 198	- 198	- 198	192 - 198 2	192 - 198	192 - 198 2	861		201	- 200	193	- 200	- 200	- 200	- 200	- 200	200	- 200	700	267	200	105	100	100	100	108	198	- 195	- 195
176	163 - 176	176	176	176	176	176	176	1.92	176	176	26	79 1	1	78	2	2%	78	78	- % - %	28	78	 8 2	× ;	 - %	- 178 1	173 1	176	174	172	176 1	- 176	- 173	-173
	251 16 251 16											254 16						_						· ·	, .			•	•				163
•	141 - 2		•	•								•		t	137 - 2	1	•		•	•	•	143 - 253	127 247					- (142 - 240		141 - 251		142 - 249
1074	1076	1077	1078·	1079	1080	1081	1082	1083	1084	1085	1086	1087	1088	1089	1090	1091	1092	1093	1094	2601	1090	1001	1000	1100	1101	1102	1103	1104	1105	1106	1107	1108	1109
1018A02	1018A05	I018A11	1018B02	I018B08	I018C04	I018D02	1018E06	I018E08	I018F04	1018G06	I018H07	I019E05	1019F06	1019G12	1020D01	1020D05	1020E10	1020G12	1020H06	1020H10	1021A11	1021501	1021011	1021E10	1021G02	1022A08	1022B01	1022R10	1022C02	1022C04	I022C08	1022D06	1022E08

																								•	Ξ			٠.					
5 DGYYDILTGYSYYGMDV (SEQ ID NO: 2135) 5 DGYYDILTGYSYYGMDV (SEO ID NO: 2135)		S DGYYDLLIGYSYYGMDV (SEQ ID NO: 2135)		•		_	DGYYDILTGYSYYGMDV (SEO ID NO:	DGYYDILTGYSYYGMDV (SEO ID NO:	ELGSSIVGATTGALDM (SEO ID NO: 28	-	ELGLSIVGATTGALDM (SEQ ID NO:	DQGRYLDL (SEQ ID NO: 2175)	3 DNYDILTGYSRRFDP (SEQ ID NO: 2942)	-					•	٠,		•				GDYDILTGYPAECFQI (SEQ ID NO: 2854)	5 GSVYDILTGTYYKSGMGV (SEQ ID NO: 2733)		BLGSSIVGATTGALDM (SEQ ID NO: 2852)		•	RYGDPFYYYYMNV (SEQ ID NO: 2755)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)
99 - 115 99 - 115	99-113	99 - 115		99 - 113	99 - 115	99-115	99 - 115	99-115	98 - 113	98 - 113	98 - 113	99 - 106	99-113	98 - 113	98 - 113	98-113	99 - 115	98-113	98 - 113	99 - 114	99 - 112	99 - 110		98 - 113		99 - 114			·	١.		99 - 112	98 - 113
50 - 66 50 - 66		90 - 00 20 - 66		. 99	99	99	50 - 66	50 - 66	49 - 65	49 - 65	49 - 65	99 - 09	99 - 09	49 - 65	49 - 65	49 - 65	20 - 66	49 - 65	49 - 65	20 - 66	99 - 09	99 - 09	49 - 65	49 - 65	50 - 65	20 - 66		1	49 - 65	49 - 65	20 - 66		49 - 65
26 - 35 26 - 35		26 - 35		26 - 35	26 - 35	26-35	26-35	26 - 35	26 - 34	26 - 34	26 - 34	26 - 35	26-35	26 - 34	26 - 34	26 - 34		26 - 34	26 - 34	26 - 35	26-35	26 - 35	26 - 34		26 - 35	26 - 35	26 - 35		26 - 34		26 - 35	26 ÷ 35	26 - 34
1 - 126 1 - 126	1 - 124	1 - 126	1 - 126	1 - 124	1 - 126	1 - 126	1 - 126	1 - 126	1 - 124	1 - 124	1 - 124	1-117	1 - 124	1 - 124	1 - 124	1 - 124	1 - 126	1 - 124	1 - 124	1 - 125	1 - 123	1 - 121	1 - 124	1 - 124	1 - 128	1 - 125	1 - 127	1 - 124	1 - 124	1 - 122	1 - 125	1 - 123	1 - 124
228 - 238 228 - 238	1	231 - 240			228 - 238	228 - 238	228 - 238	228 - 238	231 - 240	230 - 240	228 - 238	•	233 - 242	231 - 240	231 - 240	231 - 241	231 - 241	230 - 240	•	231 - 240	230 - 240	227 - 237	231 - 241	•	•	t	t	1		•	•	230 - 241	231 - 241
	187 - 193	- 198	- 195					189 - 195	192 - 198		- 195	- 190	- 198	- 198	-198	- 198	- 198	- 197	- 198	- 198		- 194	- 198	- 198	- 201	- 197	- 200	- 198	- 198	- 196	- 198	- 197	192 - 198
		- 176		-171	-173	-173	-173	_	- 176	- 175	- 173	_	- 176	- 176	- 176	-176	- 176	- 175	- 176	- 176	- 175	- 172	- 176	- 176	- 179	- 175	- 178	-176	- 176	- 174	- 176	- 175	164 - 176 19
-249 1	140 - 247 1	- 251	- 249	- 247	- 249	- 249	- 249	- 249	-251	- 251	- 249	-244	- 253	-251	251	-252 1	- 252 1	-251	- 252	- 251	-251	- 248	- 252	- 254	-255	- 249	- 254	-251	- 251	_	- 251	- 252	140 - 252 10
1110	1112	1114	1115	1116	1117	1118	1119	1120	1121	1122	1123	1124	1125	1126	1127	1128	1129	1130	1131	1132	1133	1134	1135	1136	1137	1138	1139	1140	1141	1142	1143	4	1145
I022F01 I022F04	I022F12	1023D01	1023D04	I024B04	I024D01	I024F06	1024H01	I024H07	I025A01	I025A04	I025A07	I025B01	I025B10	I025B12	1025C07	I025D11	1025E04	1025E05	I025E07	1025E10	I025F01	I025F08	1025G03	1025G08	1025H02	1026A01	1026B01	I026B06	1026C06	1026C10	1026C11	1026D09	I026E04

			•	
GYDDILTGYIMALDY (SEQ ID NO: 2821) ELGSSIVGATTGALDM (SEQ ID NO: 2852) GTGYDILTGYYMGSAFDO (SEO ID NO: 2860)			ELGSSIVGAL I GALLDM (SEQ ID NO: 2832) DNYDILTGYSRREDP (SEQ ID NO: 2942) ELGLSIVGATTGALDM (SEQ ID NO: 2174) GDYDILTGYPAECFQI (SEQ ID NO: 2854) DMYYDILTGYYTGLAFDM (SEQ ID NO: 2830) VLNYDILTGYYYGMDV (SEQ ID NO: 2832)	
99 - 113 98 - 113 98 - 113 98 - 113 98 - 113 98 - 113 98 - 113 98 - 113 99 - 116			98 - 113 99 - 113 98 - 114 99 - 116 99 - 114	
50 - 66 49 - 65 49 - 65 49 - 65 49 - 65 50 - 66	50 - 66 50 - 66 50 - 66 50 - 65 50 - 66	49 - 63 50 - 66 49 - 65 50 - 66 49 - 65	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 68 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
26 - 35 26 - 34 26 - 34 26 - 34 26 - 34 26 - 34 26 - 35			26 - 34 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	
1 - 124 1 - 124 1 - 124 1 - 124 1 - 124 1 - 124	1 - 124 1 - 124 1 - 124 1 - 124	1 - 124 1 - 124 1 - 124 1 - 124	1 - 124 1 - 124 1 - 125 1 - 127 1 - 127	1 - 132 1 - 126 1 - 127 1 - 125 1 - 127 1 - 127
230 - 240 231 - 240 231 - 240 231 - 240 237 - 245 231 - 240 231 - 240	1 1 1 1 1		231 - 240 233 - 242 229 - 239 231 - 241 229 - 239 231 - 240	
- 198 - 198 - 198 - 202 - 202 - 198 - 198	197 198 198 198	961 198 198 198 198	192 - 198 192 - 198 192 - 198 190 - 196 192 - 198	200 200 194 198 198 199 194
175 191 176 192 176 192 176 192 176 192 177 193			176 19 19 19 19 19 19 19 19 19 19 19 19 19	
			2 4 4 4 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	
140 - 251 140 - 251 140 - 251 140 - 256 140 - 251 140 - 251 140 - 251	139 - 251 140 - 251 144 - 255 140 - 251 139 - 250	138 - 249 141 - 250 140 - 251 140 - 252 144 - 252	140 - 251 140 - 253 140 - 250 141 - 252 141 - 251 141 - 251	
1146 1147 1148 1150 1151 1151	1154 1155 1156 1157 1158	1160 1161 1163 1164 1165	1160 1167 1168 1169 1170	1173 1174 1175 1176 1177 1178 1179 1180
	. 5) 10 5 5 5 11	•• ••		
1026E06 1026E09 1026F01 1026F12 1026G08 1026G11 1026G11	1026H02 1026H06 1026H10 1027A09 1027B02 1027B05	1027C08 1027D02 1027E03 1027E05 1027F04 1027F05	1027F11 1027G06 1027H03 1027H03 1028A04 1028A07	1028B10 1028C01 1028C04 1028C08 1028D04 1028D12 1028B12 1028E06

	٠.																																	•
ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) DDRRGYYDILTGYYRFGSFDI (SEQ ID NO: 2901)	DIDIGGDDS (SEQ ID NO: 2954)	VSGYNSGYFESYDMDV (SEQ ID NO: 2732)	EVRNYDLLTRSYLAGPLDN (SEQ ID NO: 2751)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	SGEPCITLACNLGG (SEQ ID NO: 2797)	DASEYYDILTGYYLATGRNWFDP (SEQ ID NO: 2888)	DPSPYYDILTGYFLPYYMDV (SEQ ID NO: 2843)	EIDDILTGYYMDV (SEQ ID NO: 2905)	MAYDILTGLVNWFDP (SEQ ID NO: 2786)	RDILTGFYDS (SEQ ID NO: 2933)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	EGSYDILTGYYVGVGRMDV (SEQ ID NO: 2171)	EVRNYDLLTRSYLAGPLDN (SEQ ID NO: 2751)	EVRNYDLLTRSYLAGPLDN (SEQ ID NO: 2751)	GYYDILTGYQSDAFDI (SEQ ID NO: 2927)	TERFGAKDVTARWGMDV (SEQ ID NO: 2874)	ENYDIL TGYYNFFDY (SEQ ID NO: 2737)	RQYDIL TGYYGGFDY (SEQ ID NO: 2958)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	RYGDPFYYYYYMNV (SEQ ID NO: 2755)	RYGDPFYYYYYMNV (SEQ ID NO: 2755)	RYGDPFYYYYYMNV (SEQ ID NO: 2755)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	ELGHREGGYWYSPYNV (SEQ ID NO: 2838)	RYGDPFYYYYYMNV (SEQ ID NO: 2755)	DPGNYDILTGYYYYYGMDV (SEQ ID NO: 2935)	SGPGWFDP (SEQ ID NO: 2870)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	RYGDPFYYYYYMNV (SEQ ID NO: 2755)	SGPGWFDP (SEQ ID NO: 2870)	RYGDPFYYYYYMNV (SEQ ID NO: 2755)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	RYGDPFYYYYMNV (SEQ ID NO: 2755)
99 - 114 99 - 119	99 - 107	99 - 114	99-117	99 - 114	102 - 115	99 - 121	99 - 118	101 - 113	98 - 112	101 - 110	99 - 114	99 - 117	99 - 117	99-117	99 - 114	99 - 115	99 - 113	99 - 113	98 - 113	99 - 112	99 - 112	99 - 112	98 - 113	99 - 114	99 - 112	101 - 119	99 - 106	98 - 113	98 - 113	99 - 112	99 - 106	99 - 112	98-113	99 - 112
50 - 66 50 - 66	99 - 09	20 - 66	99 - 05	20 - 66	52 - 69	20 - 66	20 - 66	50 - 68	50 - 65	20 - 68	20 - 66	99 - 09	99 - 09	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	49 - 65	99 - 09	99 - 09	20 - 66	49 - 65	50 - 6 6	20 - 66	50 - 68	20 - 66	49 - 65	49 - 65	99 - 09	20 - 66	20-66	49 - 65	20 - 66
26 - 35 26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 37	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 34	26 - 35	26 - 35	26 - 35	26 - 34	26 - 35	26 - 35	26 - 35	26 - 35	26 - 34	26 - 34	26 - 35	26 - 35	26 - 35	26-34	26 - 35
1 - 125 1 - 130	1 - 118	1 - 125	1 - 128	1 - 125	1 - 126	1 - 132	1 - 129	1 - 124	1 - 123	1 - 121	1 - 125	1 - 128	1 - 128	1 - 128	1 - 125	1 - 126	1 - 124	1 - 124	1 - 124	1 - 123	1 - 123	1 - 123	1 - 124	1 - 125	1 - 123	1 - 130	1-117	1 - 124	1 - 124	1 - 123	1-117	1 - 123	1 - 124	1 - 123
227 - 237 235 - 245		•		231 - 240	230 - 238	234 - 245	235 - 244	230 - 239	225 - 235	226 - 236	231 - 240	234 - 243	234 - 243		227 - 237	232 - 242	30 - 242	231 - 241	230 - 238	229 - 239	228 - 238	228 - 238	228 - 236	230 - 240	230 - 241		223 - 233	230 - 240		230 - 240	•	228 - 238		230 - 240
202	191	198	- 201	- 198	197	201	202	197	192	193	198	201	201	201	194	199	197	- 198	- 197	- 196	195	- 195	- 195	- 197	- 197	-204	- 190	- 197	- 197	- 197	- 190	- 195	- 197	- 197
172 188 - 180 196 -	169 185	176 192 -	179 195	176 192	175 191 -	179 195 -	- 961 081	175 191 -	170 186-	171 187-	176 192-	179 195	179 195-	179 195-	172 188-	177 193 -	175 191 -	176 192	175 191	174 190	173 189-	173 189	173 189	175 191	175 191	182 198	168 184	175 191	175 191	175 191	168 184	173 189		175 191
162-	156-	163 -	166-	163 -	165 -	169 -	167	162-	160	159-	163	166	166 -	166-	162-	165	163 -	163	163-	162-	163 -	163 -	163 -	165 -	162 -	169 -	156-	163 -	163 -	162 -	156 -	163 -	163 -	162 -
141 - 248 146 - 256	•	141 - 251	•	141 - 251	142 - 249	148 - 256	145 - 255	140 - 250	139 - 246	137 - 247	141 - 251	144 - 254	144 - 254	•	141 - 248	142 - 253		140 - 252	140 - 249	139 - 250	139 - 249	139 - 249	140 - 247	141 - 251	139 - 252	146 - 256	133 - 244	140 - 251	140 - 250	139 - 251	133 - 244	139 - 249		139 - 251
1182	1184	1185	1186	1187	1188	1189	1190	1191	1192	1193	1194	1195	1196	1197	1198	1199	1200	1201	1202	1203	1204	1205	1206	1207	1208	1209	1210	1211	1212	1213	1214	1215	1216	1217
I028E08 I028F06	I028F08	I028G08	1028G09	T028G10	I028H02	I028H03	1028H06	I028H09	I029A10	I029A12	I029B11	1029C08	1029E10	I029F08	I029G08	I030A02	I030A03	I030A04	I030A05	1030A09	I030A12	I030B06	I030B08	I030B10	I030C03	1030C06	1030C08	1030C09	1030C10	1030C11	I030C12	1030D07	I030D12	1030E02

		809) 72) 809) (9)	809) (813) (809)
	: 2914) 748)	NO: 29 NO: 29 NO: 29 O NO: 21 NO: 215 NO: 28	D NO: 2 D NO: 2 D NO: 2 D: 2131)
: 2174) : 2174) : 2174) : 2174) 2755) 2757) : 2174) : 2174)	(942) (942) (2755) (2755) (1D NO) (1D NO) (942) (2174)	: 2806) (SEQ ID (SEQ ID (SEQ ID (SEQ ID)	(SEQ II (SEQ II (SEQ II Q ID N(
	D NO: 2 D NO: 2 D NO: 3 D NO: 3 D NO: 2 D NO: 2	HONO HONO HONO HONO HONO SS) SS) SS) HONO HONO HONO HONO HONO HONO HONO HON	2136) CMDV CMDV D NO P D NO C 2136) C 2136) C C C C C C C C C C C C C C C C C C C
M (SEQ M (SEQ M (SEQ M (SEQ I (SEQ I (SEQ I (SEQ I (SEQ I (SEQ I (SEQ I (SEQ I (SEQ I (SEQ I	(SEQ II (SEQ II (SEQ II M (SEQ II MDV (SEQ II M (SEQ II	N (SEQ YHYYY YHYYY YHYYY NO: 29 'DV (SI YHYYY YHSAN	YHYY YHYY YHYY D NO YHYY YHYY YHYY
GALDI GALDI GALDI GALDI GALDI GALDI GALDI	REDP GALDI GALDI GALDI GYYYH REDP GALDI	A CONTROLL WDRY WDRY SEQ ID ASDVI WDRY TGHSS TGHSS	NOT (SECTION OF CONTROL OF CONTRO
VGATT VGATT VGATT VGATT VGATT VGATT VGATT VGATT	TGYSE VGATT VGATT VGATT YDILTRN TGYSE VGATT	TAYTI XKVKP XKVKP SSDS (\$ YDSSG XTOSC YDAL	JERAFI OKVKP JESYVE OKVKP JERAFI OKVKP
ELGLSIVGATTGALDM (SEQ ID NO: 2174) RYGDPFYYYYYMNV (SEQ ID NO: 2755) AGYDLLTGYPFYFDS (SEQ ID NO: 2757) ELGLSIVGATTGALDM (SEQ ID NO: 2755) ELGLSIVGATTGALDM (SEQ ID NO: 2174) ELGLSIVGATTGALDM (SEQ ID NO: 2174) RYGDPFYYYYYMNV (SEQ ID NO: 2174) RYGDPFYYYYYMNV (SEQ ID NO: 2174)	ALIGITATITATIVA (SEQ ID NO: 273) DNYDILTGYSRRFDP (SEQ ID NO: 2942) ELGLSIVGATTGALDM (SEQ ID NO: 2174) RYGDPFYYYYYMNV (SEQ ID NO: 2174) ELGLSIVGATTGALDM (SEQ ID NO: 2174) DRGGNYDILTGYYFHHGVDV (SEQ ID NO: 294 ATKSYDILTRMYYYHMDV (SEQ ID NO: 2942) BNYDILTGYSRRFDP (SEQ ID NO: 2942) ELGLSIVGATTGALDM (SEQ ID NO: 2174)	PYYDPLIAYTEQYFGN (SEQ ID NO: 2806) PYYDPLIAYTEQYFGN (SEQ ID NO: 2806) GREDTDKVKPWDRYYHYYMDV (SEQ ID NO: 2809) GLGHTDSDS (SEQ ID NO: 2959) AKGYYYDSSGASDVFDV (SEQ ID NO: 2871) GREDTDKVKPWDRYYHYYYMDV (SEQ ID NO: 2809) SSPPKWYDALTGHSSYHSAMDV (SEQ ID NO: 2159) SNPPKWYDALTGHSSYHSAMDV (SEQ ID NO: 2159)	GYDSSAFRAFDI (SEQ ID NO: 2136) GREDTDKVKPWDRYYHYYYMDV (SEQ ID NO: 2809) GYDSSAFRAFDI (SEQ ID NO: 2136) FYDTLTSYVFQYFDH (SEQ ID NO: 2137) GRKDTDKVKPWDRYYHYYYMDV (SEQ ID NO: 2813) GYDSSAFRAFDI (SEQ ID NO: 2136) GREDTDKVKPWDRYYHYYYMDV (SEQ ID NO: 2809) AATTSQKHNKYAYYFYGMDV (SEQ ID NO: 2131)
11	113	20 20 20 20 20 20 20 20 20 20 20 20 20 2	18 12 12 12 12 12 13 13 14 15 15 15 15 15 15 15 15 15 15 15 15 15
8 8 8 8 6 5 8 8 8 8		-	
49 - 65 49 - 65 49 - 65 49 - 65 50 - 66 50 - 66 49 - 65 50 - 65 65 - 65	50 - 60 - 60 - 60 - 60 - 60 - 60 - 60 -	50 - 66 - 66 - 66 - 66 - 66 - 66 - 66 -	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
26 - 34 26 - 34 26 - 34 26 - 34 26 - 37 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35
1 - 124 1 - 124 1 - 125 1 - 125 1 - 125 1 - 125 1 - 125 1 - 125	1 - 124	1 - 125 1 - 131 1 - 130 1 - 126 1 - 131 1 - 131	1 - 121 1 - 121 1 - 123 1 - 131 1 - 131
- 241 - 240 - 240 - 241 - 241 - 240 - 240 - 243 - 243		- 240 - 246 - 246 - 247 - 247 - 249	- 237 - 248 - 242 - 249 - 249 - 246 - 246
231 230 231 231 231 231 231 231 231 231 231 231	3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		
192 - 198 192 - 198 191 - 197 192 - 198 190 - 196 191 - 197 191 - 197 192 - 198	100000000000000000000000000000000000000	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	6.7793898
176 176 177 176 177 177 177 177 177 177			•
165 163 163 163 163 163 163 163 163			
140 - 252 140 - 251 140 - 251 140 - 252 140 - 252 140 - 251 140 - 253 140 - 253			
1218 1220 1221 1222 1223 1225 1226 1226	1229 1230 1231 1232 1233 1234 1235	1238 1239 1240 1241 1242 1243 1244	1246 1247 1248 1249 1250 1251 1253
E05 E07 E08 E10 E10 F05 F05 F05 F06 F11	F112 G03 G07 H06 H10 H11	A03 A03 A12 B06 B06 B07 308	811 C02 C02 C03 C04 C04 C04 C04 C07 C04
1030E05 1030E07 1030E08 1030E09 1030E02 1030F05 1030F06 1030F08 1030F08	1030512 1030503 1030503 1030509 1030609 10306110	1031A03 1031A08 1031A12 1031B06 1031B07 1031B08	1031B11 1031C01 1031C02 1031C04 1031C08 1031C11 1031D01

	_																																	
-	GREDIUKVKLWUKYYHYYYMDV (SEQ ID NO: 2807)	•		DKAHGEYGRDYYYYYGMDV (SEQ ID		SGPPKWYDALTGHSSYHSAMDV (SEQ	SSPPKWYDALTGHSSYHSAMDV	GREDTDKVKPWDRYYHYYYMD	GYDSSAFRAFDI (SEQ ID NO: 2136)				GYDSSAFRAFDI (SEQ ID NO: 2136)	DTVRSGGMDV (SEQ ID NO: 2804)	GREDIDKVKPWDRYYHYYYMDV (SEQ ID NO: 2809)	DKAHGEYGRDYYYYYGMDV (SEQ	_	GYDSSAFRAFDI (SEQ ID NO: 2136)		GREDTDKVKPWDRYYHYYYMDV (SEQ ID NO: 2809)	_	SSPPKWYDALTGDSSYHSAMGV (SEQ ID NO: 2816)	_	AATTSQKHNKYAYYFYGMDV (SEQ ID NO: 2131)	_		_	_	DKAHGEYGRDYYYYYGMDV (SEQ ID NO: 2735)	DRGYTGYDRLVGGYYFDF (SEQ ID NO: 2931)	DTVRSGGMDV (SEQ ID NO: 2804)	_		EVRNYDLLTRSYLAGPLDN (SEQ ID NO: 2751)
	99 - 120			99 - 117	99 - 120	99 - 120	99 - 120	99 - 120	99 - 110	99 - 120	99 - 120	99 - 110	99 - 110	99 - 108	99 - 120	99-117	99 - 110	99 - 110	99 - 120	99 - 120	99 - 120	99 - 120	99 - 120	99 - 118	99 - 110	99 - 115	99 - 120	99 - 117	99 - 117	99 - 116	99 - 108			99 - 117
50 - 66	20 - 66	50-66	99 - 09	20 - 66	50 - 66	20 - 66	50 - 66	20 - 66	99-09	50 - 66	99-09	20 - 66	20 - 66	50 - 66	20 - 66	99 - 09	20 - 66	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	52 - 69	52 - 69	20 - 66
26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 37	26-37	26 - 35
1-121	1 - 131	1 - 121	1 - 131	1 - 128	1-131	1-131	1-131	1-131	1 - 121	1 - 131	1 - 131	1-121	1 - 121	1-119	1 - 131	1 - 128	1-121	1 - 121	1 - 131	1-131	1-131	1-131	1 - 131	1 - 129	1 - 121	1 - 126	1 - 131	1 - 128	1 - 128	1 - 127	1 - 119	1 - 128	1 - 128	1 - 128
•	237 - 247		236 - 245	233 - 243	237 - 247	236 - 246	237 - 248	238 - 248	228 - 235	237 - 247	237 - 247	227 - 237	227 - 235	226 - 236	•		1	227 - 237	237 - 247	•		238 - 248					•	234 - 246	234 - 244		225 - 235		•	234 - 243
- 194	96 - 202	193	97 - 203	94 - 200	98 - 204		-		89 - 195	98 - 204	98 - 204		88 - 194	87 - 193					98 - 204															95 - 201
160 - 172 1	170-182 167-180	7	1-181 1	8-178 1	0-182 1	1-181 1	0-182	0-183	2-173 1	0-182	0 - 182	0-172 1	2-172 1	9-171 1	$\overline{}$	7-179 1	0-172 1	0-172 1	0 - 182	7	7	•	ī	7	0-173	7	0-183	7-179	1-179 1	•	8-170 1	6-179 1		6-179 1
	- 257 167	_	-256 171	_	_	_	_			_	_	_	_		_			•			. 258 170	_			_	_	_	_	_	_	٠.	٠.		-254 166
137	4	137.	147	144	147.	147	147	147	137	147	147	137.	137	135	147.	144	137.	137.	147	147.	147	147.	147	145	137-	142	147	144	14 4	143	135	14	144	144
1254	1256	1257	1258	1259	1260	1261	1262	1263	1264	1265	1266	1267	1268	1269	1270	1271	1272	1273	1274	1275	1276	1277	1278	1279	1280	1281	1282	1283	1284	1285	1286	1287	1288	1289
1031D04	1031D08	I031D09	1031D11	1031D12	1031E01	I031E05	I031E07	I031E08	1031E09	I031E10	I031E11	I031F01	I031F04	I031F06	I031F10	I031F11	1031F12	1031G01	I031G03	1031G05	I031G06	I031G07	I031G09	1031G12	I031H01	I031H02	1031H03	I031H06	1031H09	1031H10	I031H11	I033A08	I033B11	I033C01

											~																	=							
EMGYDILTGYYLNYMDV (SEQ ID NO: 2862)	-	·		•							Ī			•			•								DRRDYDLLTRYYYYYGMDV (SEQ ID NO. 2928)	KQRGDYDILTGYQLGYAFDI (SEQ ID NO: 2808)	SHYDILTRLNYWYFDL (SEQ ID NO: 2950)		_	Ť			SHYDILTGLNYWYFDL (SEQ ID NO: 2166)	QQWLPYDAFDI (SEQ ID NO: 2839)	AYYDILTGYFFDI (SEQ ID NO: 2873)
99 - 115	99-111	99 - 114	99 - 114	99 - 113	99 - 117	99 - 112	99 - 114	99 - 116	99-115	99 - 112	99 - 118	99 - 117	99 - 107	97 - 116	99 - 115	99 - 114	98 - 113	99 - 112	99 - 114	99-117	98 - 115	99 - 114	99 - 113	99 - 109	99 - 117	98 - 117	99 - 114	99 - 119	99 - 113	99 - 117	98 - 115	99-113	99 - 114	99 - 109	101 - 113
99 - 05	20 - 66	20 - 66	50 - 66	50 - 66	50 - 66	50 - 66	99 - 09	50 - 66	50 - 66	99 - 09	50 - 66	50 - 66	50 - 66	48 - 64	99 - 09	90 - 09	49 - 65	50 - 66	50 - 66	50 - 66	50 - 65	20 - 66	99 - 09	99 - 09	20 - 66	20 - 65	20 - 66	20 - 66	20 - 66	50 - 66	50 - 65	20 - 66	20 - 66	99 - 09	20 - 68
26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	24 - 33	26 - 35	26 - 35	25 - 34	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35	26 - 35
1 - 126	1 - 122	1 - 125	1 - 125	1 - 124	1 - 128	1 - 123	1 - 125	1 - 127	1 - 126	1 - 123	1 - 129	1 - 128	1 - 118	1 - 127	1 - 126	1 - 125	1 - 124	1 - 123	1 - 125	1 - 128	1 - 126	1 - 125	1 - 124	1 - 120	1 - 128	1 - 128	1 - 125	1 - 130	1 - 124	1 - 128	1 - 126	1 - 124	1 - 125	1-120	1 - 124
228 - 238	226 - 234	231 - 240	227 - 237	226 - 236	234 - 243	228 - 238	231 - 240	232 - 242	228 - 238	225 - 235	234 - 245	234 - 243	220 - 230	•	228 - 238		226 - 236		230 - 240	232 - 240	232 - 241	228 - 238	230 - 239	226 - 235	230 - 240	230 - 240	231 - 240	236 - 245	230 - 239			226 - 236			230 - 239
- 195	- 193	•	- 194 2	- 193	- 201	- 195	198	199	195	192	201	- 201	187	200	195	194	193	192	197	661			197	193	197		198	203	197	661	197	 83	194	•	- 197 2
189	187	192	188	187	195	189	192	193	189-	186-	195	195	181	194 -	189-	188	187	186-	191	193	193	189	191	187-	191	191	192	197	191	193	191	187-	188-	187.	191
163 - 173	-12	- 17	162 - 172	- 17	-17	- 17	163 - 176	-17	163 - 173	- 17	- 17	166 - 179	- 16	- 17	-17	- 17	-17	- 17	163 - 175	167 - 177	164 - 177	53 - 173	162 - 175	•	•	65 - 175	163 - 176	181 - 891	62 - 175	121 - 121	165 - 175	161 - 171	162 - 172	58 - 171	162 - 175
				-		•			_										_		_	_	_	_	_	_			_		_	_	_	_	_
•	138 - 245	141 - 251		•	144 - 254	139 - 249	141 - 251		142 - 249		•	144 - 254	134 - 241	•	142 - 249	141 - 248	140 - 247	139 - 246	ŧ	•	.1	•	1	•	•			•	140 - 250	٠		140 - 247	141 - 248	• •	140 - 250
1290	1291	1292	1293	1294	1295	1296	1297	1298	1299	1300	1301	1302	1303	1304	1305	1306	1307	1308	1309	1310	1311	1312	1313	1314	1315	1316	1317	1318	1319	1320	1321	1322	1323	1324	1325
1033C08	10331202	1033D03	1033D05	1033D11	1033D12	1033E01	1033E06	I033E11	I033E12	I033F03	1033F08	1033F10	I033F12	1033G01	I033G03	1033G08	1033H04	I037A05	I037B03	1037B04	1037C04	1037C06	1037C08	1037011	1037E06	1037F04	1037G01	1037G03	1037G10	I042A07	1042A10	I042B03	I042B12	I042D01	I042D03

																														_					
	EQ ID NO: 2802)	SEC ID NO: 2798)	J NO: 2883)	J NO: 2/38)	D NO: 29/0)	D NO: 2907)	D NO: 2166)	OF TO 12 (28)	(SEC ID NO: 2899)	(SEQ ID NO: 2/44)	7 NO: 2133)	54 ID INO: 2020)	3EO ID MO: 2727	FO ID NO: 2751)	O ID NO: 2940)	NO. 2153)	MO: 2153)	O IO IO IO: 2940)		TO NO. 2968)	(O. 2978)	(C. 23/6)	FO ID NO: 2776)	SEO ID NO: 2889)	NO. 2153)	2825)	O ID NO: 2758)	DV (SEO ID NO: 2912)	NO: 2153)	KN (SEO ID NO: 2845)	EO ID NO: 2751)) NO: 2946)	(D NO: 2829)	D NO: 2728)	NO: 2731)
O A TOWN TOWN TOWN OF THE PARTY	ENALTHING IF TOWN (SEQ ID NO: 2802)	CALLIDELOTIVITORNO (SEQ ID NO: 2798)	GUNDII TGVDI UARDI (SEQ ID NO: 2883)	OF THE TOYNEGAIN (SEQ ID NO: 2/38)	HVDII TGVGI I GVANV (SEO	SHYDII TGI MYINYEDI (SEO ID NO: 290)	GRI VYDII TRVVICNA FDI (SEQ ID NO: 2166)	GYYDII TEGRYYYYCHDY	GGVVDII TGVI VXVGVANV (GEO ID NO: 2899)		OOOVDII TGVHIDVAAMAY (SEQ ID NO: 2133)	ATYDPI, TGYSEDGEDI (SEO ID NO: 2152)	HVRDYDII.TGYVRGHHEDY (SEO ID NO: 2727)	EVRNYDLI TRSYI AGPI DN (SPO ID NO: 2751)	TESNYDII.TGYYWPSMDV (SEO ID NO: 2940)	ATYDPI TGYSFDGEDI (SEO 1D NO: 2153	ATYDPI.TGYSEDGEDI (SEO ID NO: 2153	TESNYDII.TGYYWPSMDV (SEO ID NO: 2	ATYDPI.TGYSFDGFDI (SEO ID NO: 2153)	APYDILTGYSDYYGMDV (SFO ID NO: 2968)	DSDARLAALDAFDI (SEO ID NO: 2978)	GOEGVI PNYVYHMOV (SEO ID NO: 29/3)	DIKRYNSNWPYYDYYMDV (SEC ID NO: 2345)	DKOYYDILTGDPVEGGMDV (SEO ID NO: 2889)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153	AGSSLVTYGTDV (SEO ID NO: 2825)	SDDYDILTGNYVGSLLDY (SEO ID NO: 2758)	DGRLSYDILTGYYARDYYGMDV (SEO ID NO	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	DONHPIYDIL TGYYVPTGPLELKN (SEO ID NO. 2845)	EVRNYDLLTRSYLAGPLDN (SEO ID NO: 2751)	DMGYDILTGYYGAFDI (SEO ID NO: 2946)	DYYDVLTGFSLDGMDV (SEQ ID NO: 2829)	DHYDVLTGSYLQAFDV (SEQ ID NO: 2728)	GRYDFLTGYLRNFDY (SEQ ID NO: 2731
08 - 115 E	_		115		2 7	7	114	2 2	. ×	. 411	- 116	- 114	- 116	117	- 116	- 114	4	. 91	14	_	- 112		911	117	114		99-116 SI	99 - 120 D	99-114 A	99 - 122 D	99-117 E	99-114 D	- 114	- 113	100 - 114 GI
59-05	52.69	20 - 66	52 - 67	20-66	20.66	20 - 66	50 - 65	20.05	50 - 65	20 - 66	20 - 66	50 - 66	50 - 65	50 - 66	99 - 09	50 - 66	. 99 - 09	20 - 66	20 - 66	89 - 09	50 - 66	20 - 66	20 - 66	99 - 09	99 - 09	99-09	99 - 09	20 - 66	99 - 09	20 - 66	20 - 66	20 - 66	99-09	50 - 65	22-67
26-35	26-37	26 - 35	26 - 37	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26-35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35		26 - 37
1 - 126	1-131	1 - 124	1 - 126	1 - 124	1 - 125	1 - 125	1 - 127	1 - 128	1 - 126	1 - 125	1 - 127	1 - 125	1-127	1 - 128	1 - 127	1 - 125	1 - 125	1 - 127	1 - 125	1 - 128	1 - 123	1 - 124	1-127	1 - 128	1 - 125	1-121	1 - 127	1-131	1 - 125	1 - 133	1 - 128	1 - 125	1 - 125	1-124	1 - 125
232 - 241		•									•		233 - 242	234 - 243	229 - 239	231 - 240	231 - 240	229 - 239	231 - 240	230 - 240	228 - 238	229 - 239	232 - 242	234 - 243	231 - 240	227 - 236		1				•		1	232 - 242
93 - 199			- 199	- 193	194		200	- 201	195	- 198		192 - 198	194 - 200 3	201	190 - 196		192 - 198	90-196	192 - 198 2	- 197	- 195	190 - 196 2	193 - 199 2	-201	861	188 - 194 2	- 700	204	198	- 206	- 197	- 199	- 199	- 197	- 199
164 - 177 1	- 182	161 - 171 1	164 - 177 1	121-191	162-172 1	192-176	165 - 178 19	166-179	163 - 173 1	- 176	165 - 178 1	163 - 176 19	165 - 178 19	- 179	164 - 174 19	- 176		-174 1	_	- 175	- 173	162 - 174 19	165 - 177 19	- 179	- 176	- 172	- 178	- 182	- 176	- 184	- 175	- 177	-177	- 175	4 - 177 193
- 252	-257	- 247	- 252	- 247	- 248	-251	-253	- 254	- 249	-251	- 253	-251	- 253	- 254	- 250	-251	-251	- 250	- 251	- 251	- 249	-250	- 253	- 254	-251	- 247	-253	-257	251	259	- 251	- 253	- 253	- 248	-253 164
326 142	327 147	328 140	329 142	1330 140	331 141	332 141	333 143	334 144	1335 142	1336 141	1337 143	•	339 143	Ξ.	141 143	_	143 141		345 141	•	•	1348 140		•	_	_	353 143	- · ·						360 140	01 141
-	-		~	-	-		H	=	H		=	=		::	:	=	13	=	13	= 1	13	13	13	EI .	Ξ:	E :	E ;	<u>.</u>	£ ;	13	S :	E :	13 13	<u>.</u>	
I042D10	1042E10	1042E11	1042F08	I042F12	.I042G08	1042G10	I042H03	I043A03	I043B02	I043B03	I043B06	I043B07	I043B09	1043D11	1043E05	I043F01	I043F04	I043F12	I043H07	1044A11	1044B11	1044009	1044C10	I044D03	1044D09	1044E07	1044E11	1044507	1044502	1044G07	1044H01	TOSOAUL	1050501	102000	102000

				•
	DTVRSGGMDV (SEQ ID NO: 2804) DGSYDILTGYYIDNYMDV (SEQ ID NO: 2154) SGPGWFDP (SEQ ID NO: 2870) SGPGWFDP (SEQ ID NO: 2870) ELGSSIVGATTGALDM (SEQ ID NO: 2852) GDYDILTGYPAECFQI (SEQ ID NO: 2854) DNYDILTGYSRRFDP (SEQ ID NO: 2942)	ELGLSIVGATTGALDM (SEQ ID NO: 2174) ELGLSIVGATTGALDM (SEQ ID NO: 2174) ELGLSIVGATTGALDM (SEQ ID NO: 2174) EGGNYDELTGYYIGNGAFDI (SEQ ID NO: 2158) ATYDPLTGYYIGNGAFDI (SEQ ID NO: 2153) TYYDPLTGYYFDY (SEQ ID NO: 2153)		GGELVWFGESDYYGMDV (SEQ ID NO: 2787) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) SQRLFIDS (SEQ ID NO: 2842) DRYYDILTGYYIPGLDDAFDI (SEQ ID NO: 2887) DSDARLAALDAFDI (SEQ ID NO: 2978) EESYYDILTGYYVHYYGMDV (SEQ ID NO: 2743) ATYDPLTGYYFDGFDI (SEQ ID NO: 2949) ATYDPLTGYYFDGFDI (SEQ ID NO: 2153) AYYDILTGFLFYDMDL (SEQ ID NO: 2171)
	99 - 108 99 - 116 99 - 106 99 - 107 99 - 114	98 - 113 98 - 113 98 - 113 97 - 116 99 - 114		99-115 99-114 99-106 99-119 99-114 99-114
50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 51 - 67	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	49 - 65 49 - 65 49 - 65 50 - 66	50 - 68 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 6
26-35 26-35 26-35 26-35 26-35 26-35 26-35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35		26-35 26-35 26-35 26-35 26-35 26-35 26-35	26 - 35 26 - 35
1-124 1-121 1-125 1-131 1-131 1-127	1 - 119 1 - 127 1 - 117 1 - 117 1 - 124 1 - 125	1 - 124 1 - 124 1 - 124 1 - 127 1 - 125	1-125 1-124 1-125 1-125 1-125 1-125	1-126 1-125 1-117 1-130 1-129 1-125 1-125 1-125
	225 - 235 229 - 239 224 - 233 224 - 233 231 - 240 230 - 240 233 - 242			230 - 240 231 - 240 219 - 229 236 - 245 225 - 235 231 - 240 231 - 240 231 - 240
198 199 205 198 199	192 193 193 193 193	198 208 20	2 4 8 6 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	197 198 198 198 198
75 192- 77 193- 78 199- 76 192- 77 193-	170 186 - 174 190 - 169 185 - 169 185 - 176 192 - 176 192 -			175 191 - 176 192 - 164 180 - 170 186 - 176 192 - 176 192 - 176 192 - 176 192 -
163 - 17 160 - 17 170 - 18 164 - 17		164 - 17 163 - 17 163 - 17 163 - 17		163 - 17 163 - 17 154 - 10 160 - 17 166 - 17 163 - 17 163 - 17
	135 - 246 143 - 250 133 - 244 133 - 244 140 - 251 140 - 253	140 - 252 140 - 249 140 - 252 143 - 253 141 - 251		142 - 251 141 - 251 133 - 240 146 - 256 139 - 246 145 - 252 141 - 251 141 - 251
1362 1363 1364 1365 1366 1366	1369 1370 1372 1373 1374	1376 1377 1378 1379 1380	1382 1383 1384 1385 1386 1387	1389 1390 1391 1392 1393 1394 1395 1396
1050E01 1050E10 1050H08 1051A04 1051A08 1051A12 1051B08	1051C06 1051G12 1055A05 1055A11 1061A03 1061A04	1061A09 1061A10 1061B07 1061B09 1061B12	1061D01 1061D03 1061D04 1061D07 1061D10 1061D10	1061E05 1061E09 1061E12 1061F01 1061F10 1061F11 1061G01

	٠.																																	
28 26-35 50-66 99-117 EVRNYDLLTRSYLAGPLDN (SEQ ID NO: 2751) 27 26-35 50-65 98-116 EGSYDILTGYYYGYGRMDV (SEQ ID NO: 2171)	26-35 50-68 101-110	26-37 52-67	126 26-35 50-68 101-115 DFYDILTGYQHGMDV (SEQ ID NO: 2919)	26-35 50-66	26-35 50-66 99-111	26-35 50-66 99-117	26-35 50-66 99-106 1	129 26-35 50-66 99-118 VNADYDILTGYPRDYYGMDV (SEQ ID NO: 2819)	25 26-35 50-66 99-114 ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	26-35 50-66 99-119	26-35 50-66 99-116 1	26-35 50-66 99-113 1	26-35 50-66 99-117 1	124 26-35 50-65 98-113 GERDILTGYYLDGMDV (SEQ ID NO: 2948)	26-35 50-66 99-113 1	26-35 50-66 99-115]	26-35 50-66 99-118 1	26-35 50-66 99-114	26-35 50-66 99-	26-35 50-66 99-112	26-35 50-66 99-113	26-35 50-66 99-	26-35 50-66 99-	26-35 50-66 99-110	26-35 50-66 99-108	26-35 50-66 99-115	26-35 50-66 99-112	26-35 50-66 99-117	26-35 50-66	26-35 50-66 99-110	26 - 35 50 - 66 99 - 108	26-35 50-66 99-115 1	26-35 50-66 99-116 1	119 26-35 50-66 99-108 GMGDHYGMDV (SEQ ID NO: 2161)
254 166-179 195-201 234-243 1-128 253 165-178 194-200 233-242 1-127	159 - 171 187 - 193 226 - 236 1	-177 193 - 199 232 - 241 1 -	249 163 - 173 189 - 195 228 - 238 1 - 12	-173 189-195 228-237 1-	160 - 173 189 - 195 228 - 237 1 - 1	166-179 195-201 234-243 1-	155 - 168	-180 196-202 235-244 1-	251 163 - 176 192 - 198 231 - 240 1 - 125	_	198	162 - 174 190 - 196 229 - 240 1 -	-179 195-201 234-243 1-	197 230 - 239 1 -	196 229 - 239 1	164 - 177 193 - 199 232 - 241 1	198 231 - 241 1	248 162 - 172 188 - 194 227 - 237 1 - 125	-178 194-200 233-242 1	161 - 173 189 - 195 228 - 238 1	250 162 - 174 190 - 196 229 : 239 1 - 124	-174 190-196 229-237 1	-169 185 - 191 224 - 234 1	159-172 188-194 227-236 1	156 - 166 182 - 188 221 - 231 1	163 - 173 189 - 195 228 - 238 1	160-170 186-192 225-235 1-	179 195-201 234-243 1-	-201 234 - 243 1 -	159-171 187-193 226-236 1-	157 - 169 185 - 191 224 - 234 1	164-177 193-199 232-241 1	165-178 194-200 233-242 1-	242 156 - 166 182 - 188 221 - 231 1 - 11
144-	137-	1401 142-	•	1403 138 -	138-	1405 144 -	133	145 -	•	146 -	143 -	1411 140 -	144	1064E07 1413 140-25	1414 140-	1415 142-	145-	1064G06 1417 141 - 24	143 -	1419 139-	1065C09 1420 140 - 25	1421 141 -	1422 135 -	1423 137	1424 135	1425 142 -	1426 139-	144 -	1066A03 1428 144-25	137 -	135 -	1431 142 -	1432 143 -	135-

AGSSLMTYGTDV (SEO ID NO: 2773)	GLYFEDTNYRHGDAFDI (SEO ID NO: 2790)	GMGDHYGMDV (SEO ID NO: 2161)	ATYDPI.TGYSEDGEDI (SEO ID NO: 2153)	GMGDHYGMDV (SEO ID NO: 2161)	ENYDELTGYYGAEDI (SEO ID NO: 2772)	HSKEYNWNYAI DY (SEO ID NO: 2754)	ERSOFDFLTGVDRYHPMDV (SEO ID NO. 2956)	EGAADYI.NGOYFOH (SRO ID NO: 2815)	AGSSI.MTYGTDV (SEO ID NO: 2773)	GMGDHYGMDV (SEO ID NO: 2161)	GLYFEDTNYRHGDAFDI (SEO ID NO: 2790)	VYYDILTGHPTYGMDV (SEO ID NO: 2791)	GIYDILTGYHWDDAFDI (SEO ID NO: 2872)	ESTYDIL TGSYHDYGLDV (SEO ID NO: 2822)	DRLHYDILTGHOTDDAFDI (SEO ID NO: 2885)	VLTNYDILTGYYREDAFDM (SEO ID NO: 2939)	GMGDHYGMDV (SEO ID NO: 2161)	DRGASNYDILTGYYAPAOGVAFDI (SEO ID NO: 2969)	EGAHYDIL TGHNYYHYGMDV (SEO ID NO: 2747)	ETRKYTSSPPYNYYYMDV (SEO ID NO: 2736)	AGSSLMTYGTDV (SEO ID NO: 2773)	DQHDILTGVYYGMDV (SEQ ID NO; 2921)	GMGDHYGMDV (SEQ ID NO: 2161)	DYPGSEYDILTGYLFGYYYYGMDV (SEQ ID NO: 2926)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ARRVGVLGGKNAFEI (SEQ ID NO: 2765)	DQHDILTGGYYGMDV (SEQ ID NO: 2894)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	EGTYYDILTGYYPLGYFDY (SEQ ID NO: 2936)	GMGDHYGMDV (SEQ ID NO: 2161)	GGSSONFYGMDV (SEO ID NO: 2884)	GTGYDILTGYYMGSAFDO (SEO ID NO: 2800)	GVVWVAYGDVGIYGFDV (SEO ID NO: 2937)	HDYYIMTAAHYYYDS (SEO ID NO: 2909)	GIGYDLLTGYFTGSPLDY (SEQ ID NO: 2846)
99 - 110	99 - 115				99-113				99-110			99 - 114	101 - 117	99 - 116	98 - 116	99 - 117	99 - 108	99 - 122	102 - 121	99 - 116	99 - 110	99 - 113	99 - 108		99 - 114	99 - 113		99 - 114	101 - 119	99 - 108	99 - 110	99 - 116	99-115	99 - 113	99 - 116
50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	50 - 68	50 - 66	50 - 65	50 - 66	20 - 66	50 - 66	52 - 69	50 - 66	20 - 66	50 - 66	20 - 66	50 - 67	20 - 66	20 - 66	20 - 66	20 - 66	20 - 68	20 - 66	20 - 66	50 - 66	50 - 66	50 - 66	20 - 66
26 - 35	26 - 35	26 - 35				26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 37	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26-35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35
1 - 121	1 - 126	1-119	1 - 125	1 - 119	1 - 124	1 - 122	1 - 128	1 - 123	1 - 121	1 - 119	1 - 126	1 - 125	1 - 128	1 - 127	1-127	1 - 128	1-119	1 - 133	1 - 132	1 - 127	1-121	1 - 124	1 - 119	1 - 134	1 - 125	1 - 124	1 - 124	1 - 125	1 - 130	1-119	1-121	1 - 127	1 - 126	1 - 124	1 - 127
227 - 236	232 - 241	224 - 234	231 - 240	221 - 231	230 - 239	228 - 237	234 - 243	229 - 238	226 - 236	221 - 231	228 - 238	229 - 237	233 - 243	232 - 243	233 - 242		225 - 234	239 - 248	238 - 247	233 - 242		229 - 238		1				•		224 - 234	227 - 237	233 - 243	233 - 241	229 - 240	233 - 243
188 - 194	193 - 199	185 - 191	192 - 198	182 - 188	191 - 197	189 - 195	95 - 201	90 - 196	87 - 193	82 - 188	89 - 195	90 - 196	94 - 200	93 - 199	94 - 200	95 - 201	86 - 192	00 - 206	99 - 205	94 - 200			85 - 191	00 - 206	92 - 198	91 - 197	90 - 196	92 - 198	•	-8		94 - 200	94 - 200	90 - 196	94 - 200
59 - 172	•	157 - 169	163 - 176	156 - 166	162 - 175	160 - 173	166 - 179 1	161 - 174	129 - 171 1	56 - 166 1	163 - 173 1	164 - 174 1	166 - 178 1	165 - 177 1	165 - 178 1	166 - 179 1	57-170 1	71 - 184 2	70 - 183	165 - 178 1	7	7	7	7	163 - 176	162 - 175 1	7	7	•	57 - 169 1	160 - 172	166 - 178	65-178 1	164 - 174	166 - 178
- 247	- 252	- 245	- 251	135 - 242 1	- 250	138 - 248 1	- 254	139 - 249 1	- 247	- 242	142 - 249 1		- 254	- 254	- 253	- 254	- 245	- 259 1	- 258	- 253	- 247	- 248	-245	- 260	-251	- 250	- 250	- 252	- 256	- 245	- 248	- 254	- 252	- 251	143 - 254
		•	•	_	1439	1440	1441	1442	1443	7		_	7		_	_	_	_	_	_	_	_	_	1458 1	- ·	- '	1461 1	_	_	-	_	_			1469 1
I066B08	1066B10	I066C02	I066C11	I066C12	1066D06	1066D08	I066Di1	I066D12	1066E06	1066E12	1066G05	1066G08	1066G10	1066G12	1066H04	1067A07	I067A11	1067B08	1067C08	1067C09	1067D07	1067E01	1067E06	1067E07	1067E11	106/503	106/G05	106/G12	1067H05	1067H06	1068C09	1068G03	1068G04	1068G07	1068G08

																	•										_								
DEYDIL TGYHDAFDI (SEO ID NO. 2910)			AAYDPLTGYSFDGFDI (SEO ID NO: 2783)	DMHYDILTGYYTGLAFDM (SEO ID NO: 2917)	_		SSNPVYGLDV (SEO ID NO. 2957)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	DDRDILTNYYLEYFOH (SEO ID NO. 2868)	SSPPKWYDALTGDSSYHSAMDV (SFO ID NO: 2165)	DKTLGDOLVEAYYYDGMDV (SEO ID NO: 2776)	LGRTSRDLLTGYHFYNMDV (SEO ID NO: 2944)	DDYDILTGSLYYFDS (SEO ID NO: 2803)	GTGYDILTGYYMGSAFDO (SEO ID NO: 2800)	DRADILTGYNDAFDI (SEO ID NO: 2739)	RYGDPFYYYYMNV (SEO ID NO: 2755)	GTGYDILTGYYMGSAFDO (SEO ID NO: 2800)	VSNDILTGWGGYNWFDP (SEO ID NO: 2955)	GTGYDILTGYYMGSAFDO (SEO ID NO: 2800)	DOGRYLDL (SEO ID NO: 2175)	DOGRYLDL (SEO ID NO: 2175)	ELGLSIVGATTGALDM (SEO ID NO: 2174)	GTGYDILTGYYMGSAFDQ (SEQ ID NO: 2800)	TYYDIL TGYYAEYFQH (SEQ ID NO: 2932)	SDYDILTGYYWVPAV (SEQ ID NO: 2812)	GREDTDKVKPWDRYFHYYYYMDV (SEQ ID NO: 2835)	DQGRYLDL (SEQ ID NO: 2175)	GTGYDILTGYYMGSVFDP (SEO ID NO: 2897)	SYYDILTGYYHTPLDY (SEQ ID NO: 2853)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	GTGYDILTGYYMGSAFDQ (SEQ ID NO: 2800)	GGGYDILTGYSYPYLYYGLDV (SEQ ID NO: 2865)	GRGYDVLTGYFTGSPLDY (SEQ ID NO: 2881)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)
99 - 113	101 - 113	99 - 114	99 - 114	99 - 116	100 - 115	99 - 111	99 - 108	99 - 114	99 - 114	99 - 114	99 - 120		99 - 117	99-113	99-116	99 - 113	99 - 112	99 - 116		99 - 116	99 - 106	90 - 66	98 - 113	99 - 116	99 - 114	99 - 113	99 - 120	90 - 66	99-116		98 - 113	99 - 116	101 - 121		98 - 113
50 - 66	50 - 68	50 - 66	50 - 66	20 - 66	51 - 67	50 - 66	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	50 - 66	50 - 66	50 - 66	50-66	50 - 66	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	20 - 66	49 - 65	50 - 66	20 - 66	20 - 66	20 - 66	50 - 66	50 - 66	20 - 66	49 - 65	20 - 66	52 - 68	20 - 66	49 - 62
26-35	26-35	26-35	26 - 35	26 - 35	27 - 36	26 - 35	26-35	26-35	26-35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 34	26-35	26-35	26 - 35	26 - 35	26 - 35	26-35	26-35			28 - 37		26 - 34
1 - 124	1 - 124	1 - 125	1 - 125	1 - 127	1 - 126	1 - 122	1 - 119	1 - 125	1 - 125	1 - 125	1-131	1 - 128	1 - 128	1 - 124	1 - 127	1 - 124	1 - 123	1 - 127	1 - 126	1 - 127	1-117	1-117	1 - 124	1 - 127	1 - 125	1 - 124	1-131	1-117	1 - 127	1 - 125	1 - 124	1 - 127	1 - 132	1 - 127	1 - 124
226 - 236	230 - 239	227 - 237	231 - 240	- 1	231 - 241	228 - 237	224 - 234	231 - 240	231 - 240	229 - 239	237 - 248	233 - 242	233 - 244	229 - 239	240 - 248	229 - 239	230 - 240	232 - 242	233 - 243		223 - 233	224 - 236	230 - 241	234 - 246	•	•			234 - 242			•			231 - 241
- 193	91 - 197	188 - 194	92 - 198	90 - 196	92 - 198	89 - 195	- 191	- 198	- 198	- 196	198 - 204	194 - 200		- 196	- 205	- 196	197	193 - 199	194-200	- 199	_	- 191	197	- 201	198	- 198	- 202	- 190	- 201	- 198	- 197	- 199	- 206	-201	- 198
-171 187	- 175 19	- 172 18	176 19	- 174 19	176 19	173 18	169 185	176 192	176 192	174 190	182 19	178 19	178 19	174 190	179 195	174 19	175 191	177. 19	178 19		168 18	169 185	175 191	_					179 195	176 192	175 191	177 193	184 200		176 192
161	162	162	163			160	157.	163	163	_	170	_	168	164	166	164	162	167	165	167	156	156-	_	_	<u>2</u>	163	170	156-	168 -	164	163	167	172	166	163
140 - 247	140 - 250	141 - 248	141 - 25	143 - 250	142 - 252	138 - 248	135 - 245	141 - 251	141 - 251	141 - 250	147 - 259	144 - 253	144 - 255	•	143 - 259	1	139 - 251	143 - 253	142 - 254	ŧ	133 - 244	```		ï		•		•	``	•	ï		148 - 261	•	140 - 252
1470	1471	1472	1473	1474	1475	1476	1477	1478	1479	1480	1481	1482	1483	1484	1485	1486	1487	1488	1489	1490	1491	1492	1493	1494	1495	1496	1497	1498	1499	1500	. 1501	1502	1503	1504	1505
1070F07	1070G05	1070H02	I071A01	I071A03	I071B08	1071E01	I074F11	· I071G11	1071H08	1074A02	I074A08	I074D10	I074E01	1074E02	I074E08	I074F12	I074H06	1074H07	I074H08	I075A07	I075B01	I075B04	I075B06	I075B08	I075B09	1075B12	1075C01	I075C05	I075D05	1075/07	1075D08	1075E01	I075E03	1075E04	1075E05

ELGLSIVGATTGALDM (SEQ ID NO: 2174) SGPGWFDP (SEQ ID NO: 2870) TDRFGAKDVTARWGMDV (SEQ ID NO: 2979) EQGYDILTGYYPEGGWFDP (SEQ ID NO: 2834) AGYDLLTGYPFYFDS (SEQ ID NO: 2757) GRNYYDFLTGYNFNLGLDY (SEQ ID NO: 2830) ENYDSLTGYYNYFDY (SEQ ID NO: 2971)	DQRKAQDI (SEQ ID NO: 2779) LKAPYYDLLTGYHLPKWFDT (SEQ ID NO: 2953) DQGRYLDL (SEQ ID NO: 2175) DQGRYLDL (SEQ ID NO: 2175) BLGLSIVGATTGALDM (SEQ ID NO: 2174) GRYYDMLTRGGYFDY (SEQ ID NO: 2858) RQYDILTGYYGGFDY (SEQ ID NO: 2958) TDYDILTGYPMGYFDP (SEQ ID NO: 2173) DQGRYLDL (SEQ ID NO: 2173)	DQGRYLDL (SEQ ID NO: 2175) DQGRYLDL (SEQ ID NO: 2175) DQGRYLDL (SEQ ID NO: 2175) GSGYDLLTGYFTGSPLDY (SEQ ID NO: 2766) DRRRDDLTGYLYDAFDS (SEQ ID NO: 2878) GYDTAMQY (SEQ ID NO: 2951) DQGRYLDL (SEQ ID NO: 2175) DRRDILTGSNFGQD (SEQ ID NO: 2913) MGHYDLLTGYFTGSPLDY (SEQ ID NO: 2831) GSGYDLLTGYFTGSPLDY (SEQ ID NO: 2766)	DQGKYLDL (SEQ ID NO: 2175) DQGRYLDL (SEQ ID NO: 2175) PYYDPLTAYTFQYFGN (SEQ ID NO: 2806) ELGLSIVGATTGALDM (SEQ ID NO: 2174) GRYYDMLTRGGYFDY (SEQ ID NO: 2174) GRYYDMLTGYYPSGFDY (SEQ ID NO: 2799) RFYDLLTGYSAFDS (SEQ ID NO: 2799) GTGYDLTGYYMGSAFDQ (SEQ ID NO: 2780) GTGYDLTGYYMGSAFDQ (SEQ ID NO: 2174) GTGYDLTGYYMGSAFDQ (SEQ ID NO: 2174) GTGYDLTGYYMGSAFDQ (SEQ ID NO: 2174)
			_
98 - 113 99 - 106 99 - 115 99 - 117 100 - 114 99 - 117	99 - 106 99 - 118 99 - 106 99 - 106 99 - 113 99 - 114 99 - 116	8 8 8 9 9 8 8 8 8 8	99 - 100 99 - 110 99 - 111 99 - 113 99 - 114 100 - 113 99 - 116 99 - 116
49 - 65 50 - 66 50 - 66 50 - 66 52 - 67 50 - 66	50 - 66 50 - 66 50 - 65 50 - 65 50 - 65 50 - 66 50 - 66 50 - 66		50 - 66 50 - 66
26 - 34 26 - 35 26 - 35 26 - 35 26 - 37 26 - 35 26 - 35	26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 26 26 26 26 26 26 26 26 26 26 26 26	26 - 35 26 - 35
1 - 124 1 - 117 1 - 126 1 - 128 1 - 128 1 - 128	1 - 117 1 - 129 1 - 124 1 - 124 1 - 124 1 - 125 1 - 125 1 - 125		
231 - 241 223 - 233 233 - 243 233 - 242 231 - 240 233 - 243 230 - 240	223 - 233 236 - 246 222 - 232 224 - 234 230 - 241 231 - 241 232 - 242 233 - 243	1 1 1 1 1 1 1 1 1	224 - 234 222 - 232 - 232 239 - 239 230 - 240 231 - 241 234 - 244 229 - 239
	184 - 190 197 - 203 183 - 189 185 - 191 191 - 197 192 - 198 193 - 199 185 - 191	185 - 191 184 - 190 195 - 201 192 - 198 187 - 193 184 - 190 190 - 196 192 - 198	
- 176 - 168 - 178 - 178 - 176 - 176	156 - 168 169 - 181 157 - 167 156 - 169 163 - 175 163 - 176 164 - 177 156 - 169	- 168 - 179 - 176 - 171 - 171 - 174 - 176 - 179	827428847
-252 -244 -254 -253 -251 -251 -251	252 253 253 253 254 255 255 255 255 255 255 255 255 255		252 253 253 253 253 253 253 253 253 253
1506 1507 1508 1509 1510 1511	1513 1514 1515 1516 1517 1518 1519 1520 1521	1523 1524 1525 1526 1527 1529 1530 1531	1533 1533 1535 1536 1536 1539 1540 1541
1075E10 1075E11 1075E12 1075F02 1075F04 1075F06	1075F08 1075F09 1075F10 1075F11 1075G05 1075G07 1075G11 1075G12	1075H03 1075H06 1075H08 1076A01 1076A03 1076A07 1076A08 1076B03	1076B03 1076B04 1076C04 1076C10 1076C10 1076D11 1076D11

						٠.																		<u>@</u>										
•	·									(69)	•													DRGAPNYDIL TGYYAPAQGVAFDI (SBQ ID NO: 2176)									188)	
	6	`6							6	0.2					<u> </u>	· (C	-		•					Ö		•							2	
∃ 8	280	: 280	2			7		28)	280	DX	.	33	2	<u>ر</u>	2766	2766			6	4	4	4	4	8			4	74	74)	Æ	4	4	2	
284	2	ÖZ	: 2742)	•	2730	21.): 28	0N	350	: 297	: 286): 21	282	ö	ö	2177	2177	0:21): 21.): 21): 21): 21′	(SE		•): 21	. 21,	21,	5.21): 21,	SEQ	
N C		日 O	Š		ö	NO		NC	日夕	<u></u>	8	8	S	Ö	9	(SEQ ID NO: 2766)	ö	ö	ž	N	ž	SNO	Z	ED!			S	S	SNC ONC	N N	S S	ž		_
H OS	SE (SE	SE (SE	E C	175)	A	80	(21)	00 10 11	SE)	GMI	(SEQ ID NO: 2971	S	S	(SEQ ID NO: 2827)	(SE	(SEC	9	A	(A)	8	80	000	800	Z	(271)	(271)	80	8	80	없	8	口 ()	₩ 25	178)
V (SE	E O	FDC	Z (SE	Š	(SE	K (S)	₹Ö: 7	K (S)	PO	7	SE.	K (Si	K (S)	'(SE	20	ED ED	(SE	(SE	S) <u>X</u>	K (S)	K (S)	M (SI	K (S)	PAQ	Ş	Ö	ĭS) ¥	S) ¥	M (S)	<u>S</u>	<u>x</u>	N (S)	2	Š
EYYDVLTGLFYYMDV (SEQ ID NO: 2841) DDRDII TNYYI EYFOH (SEQ ID NO: 2868	GTGYDILTGYYMGSAFDO (SEO ID NO: 2800)	GTGYDILTGYYMGSAFDQ (SEQ ID NO: 2800)	3DYDVLTGYLRKLDY (SEQ ID NO:	DQGRYLDL (SEQ ID NO: 21 <i>75</i>)	/HYDILTGYLWAFDI (SEQ ID NO: 2730)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	OQGRYLDL (SEQ ID NO: 2175)	GRYYDMLTRGGYFDY (SEQ ID NO: 2858)	GTGYDILTGYYMGSAFDQ (SEQ ID NO: 2800)	NGYYDILTGYYLWDYYYGMDV (SEQ ID NO: 2769)	ENYDSLTGYYNYFDY	THYDILTGYYSHPLDY (SEQ ID NO: 2863)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	VPYDILTGYWGAFDV	GSGYDLLTGYFTGSPLDY (SEQ ID NO: 2766)	GSGYDLLTGYFTGSPLDY	VYYDILTGYNLFFDY (SEQ ID NO: 2177	VYYDIL TGYNL FFDY (SEQ ID NO: 2177)	MEYDILTGYYGGYFDY (SEQ ID NO: 2179)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	XX	OGGRYLDL (SEQ ID NO: 2175)	OQGRYLDL (SEQ ID NO: 2175)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	DGRLSYDILTGYYARDYYGMDD (SEQ ID NO: 2188)	SEGTIFGVD (SEQ ID NO: 2178)								
FYY.	<u> </u>	XX	(LR	(SEQ	LW	TG/	(SEQ	RGG	23	77	Z	YSH	Ę	WGA	YFT	XFI	ZLF	NLF	YGG	Ę,	βE	TG/	TG/	LTG	OES)	(SEQ	Ę	TIG/	<u> 1</u>	ğ	<u>3</u>	TTG/	167	(SEC
15.5	LTG	LTG	JG	겁	TGY	/GA7	DI,	加	LTG	LTG	TGY	IGY	'GA'	IGY	LTG	LTG	TGY	TGY	TGY	/GA]	/GA/	'GA'	/GA7	Ē	ď	ď	/GA7	/GAT	'GA'	/GA]	'GA'	'GA'		Š
DVI.	IQ.	X	TA C	RY	DIL	LSIV	RYI	Ŕ	IdX	3	DSL	口口	LSI	걸	Z	Ä	6	9	OIL TIC	LSIV	LSI	LSIV	LSIV	APN	i RYI	iryi	LSI	LSI	LSI	LSI	LSI	LSI	TSX	TEC
EYY	GIG	GTG	GDX	ğ	XHX	ELG	8	GRY	GTG	NGX	ENA	H	ELG	4	GSG	GSG	ξ	5	ME	ELG	ELG	ELG	ELG	DRG	8	8	ELG	ELG	ELG	ELG	ELG	ELG	DG	SEG
113	116	116	113	106	113	113	901	113	116	119	113	114	113	133	911	116	113	113	114	1.14	114	114	114	122	901	901	113	113	113	113	114	113	120	107
66		- 66	- 66	- 66	- 66	- 86	- 66	- 66	- 66	- 66	- 66	- 66	- 86	- 66	- 66	- 66	- 66	- 66	- 66	- 66	- 66	66	- 66	- 66		66	- 86	- 86	- 86	98 - 1	- 66	- 86	66	- 66
					•	65	 99	99	99	92	9	. 9	55	. 99	9	9	99	99	99	9,	9	9	9,	99	9	92	55	:					99	99
50 - 66	50 - 66	50 - 66	51 - 66	99 - 09	51 - 66	49 - (20-(50-(50-(50 - 66	50 - 66	99:05	49 - 65	20-(50 - 66	30-	50-(50-(50 - (20-(50-(50-0	50 - 66	50 - 66	50 - 66	50 - 66	49 - 65	49 - 65	49 - 65	49 - 65	1	49-1	20-0	20 - (
35	35	35	36	35	36	34	35	35	35	35	35	35	34	35	35	35	35	35	35	35	35	35	35	35	35	35	34	34					32	32
26-	26-	- 92	26 -	56 -	- 9 2	76-	26-	- 92	- 92	26 -	26-	26-	26-	26-	26 -	56 -	76-	76	76-	26 -	26-	5 6-	56 -	<u> 26</u> -	26-	56 -	56 -	26-	26 -	5 8	- 92	76	5 6-	5 6
- 124	- 127	127	124	117	124	124	117	124	127	130	124	125	124	124	127	127	- 124	- 124	125	125	125	125	125	133	117	117	124	- 124	124	- 124	- 125	124	- 131	118
			<u>:</u>	<u>-</u>	-	-	-	-	-	-	-	-	-	-	-		_	-	-	<u>-</u>	-	-	<u>:</u>	-	-	-	_	_	<u>-</u>	-	~	-	-	<u>-</u>
- 240		- 242	-240	-234	-239	-241	- 234	-240	- 248	- 246	- 240	- 240	-240	-241	- 245	- 245	- 239	-239	- 240	- 239	-239	-239	- 239	- 248	- 233	-233	- 238	-238			1	- 241	- 246	- 230
230 -	234 -	232	230	224	229	230 -	223	230	240 -	236 -	230	230	230 -	231	234 -	234	230.	230.	231	229	229	231	231	238	224 -	223	228	230	230	230	229	231	237	. 222
91 - 197	201	94 - 199	197	191	196	197	190	197	205	203	197	197	197	198	201	201	197	197	198	196	196	198	198	99 - 205	191	190	89 - 195	91 - 197	91 - 197	.197	196	. 198	204	. 189
191 -	195	194	191 - 1	185	190-	191-	184	191-	195	197	191	191 -	191	192-	195-	195-	191-	191.	192	190-	190-	192 -	192-	199	185-	184-	189	191	191	191 -	190-	192	198	183
175	179	178	175	169	174	175	168	175	179	181	175	175	175	176	179	179	175.	175	176	174	174	176	176	183	169	168	173	175	175	175	174	176	182	167
163 - 175	- 991	- 991	163-	157-	164-	163-	156-	163 -	- 991	- 691	163 -	165-	63-	163 -	- 991	- 991	162-	- 791	163-	164-	164-	164-	49	171	156-	156-	163-	<u>163</u> -	163 -	163 -	164-	163 -	- 69	157 -
	254	253 1	251	245 1	250 1	252	245	251	259	257]	251 1	251 1	251	252	256	256	250	250	251	250 1	250]	250	250 1	259	244	244	249	249	249	- 249	20	252	257	241
140 - 251	43 - 2	143 - 2		133-2	140-2	140-2	33-2	140-2	143-2	146-2	•	141 -2	140-2	140-2	143 - 2	143 - 2	140-2	140-2	141-2	41 - 2	41 - 2	41-2	41-2			33-2	140-2	140-2	140-2	140-2	. 1			134-2
~ ~		<u>~</u>	÷	=	~	-	=	~	_	7	7	-	7	÷	~	~	~	~	-	~	~	~	<u>~</u>	~	=	=	-	Ť	~	-	÷	<u>~</u>	~	- -
1542 1543	1544	1545	1546	1547	1548	1549	1550	1551	1552	1553	1554	1555	1556	1557	1558	1559	1560	1561	1562	1563	1564	1565	1566	1567	1568	1569	1570	1571	1572	1573	1574	1575	1576	1577
C 8	=		<u>ي</u>	¥	<u></u>	0	8	2	==		23	\$	S	90	8	2	90	¥	2	91-K	01-R	24.K	24-R	2	9	2	90	. 90	6	2	12	2	8	5
1076E07	1076E1	[076F0]	1076F03	1076F04	1076F08	1076F10	[076G09	1076G10	1076G1	1076G12	1076H02	1076H04	1076H05	1076H06	1076H09	1076H10	10777D06	[078B04	1078E10	1002A01-K	1002A01-R	026C04-K	1026C04-R	1067B10	1068C06	1075F12	1003C06	[025B06	[025B09	026C04	1027B12	1030A10	1064C04	1064C07
22	2	2	2	2	2	2	2	\approx	2	2	2	2	2	2	2	2	2	2	H	2	2	2	2	2	2	×	×	×	×	×	×	=	≍.	×

																	. ;	014)					•				•							•	
GKGYYDILTGYYRDNWFDP (SEQ ID NO: 2181)	TPSSVYDLLIGYYHYFYSYMDV (SEQ ID NO: 2189)	EKSAAGYFDY (SEQ ID NO: 2190)	ENYDSLTGYYGAFDI (SEQ ID NO: 2185)	DQGRYLDL (SEQ ID NO: 2175)	KLGLSIVGATTGALDM (SEQ ID NO: 2186)	EGMNDFINSHHYYTIMDA (SEQ ID NO: 2182)	AGNEYGHTERPADY (SEQ ID NO: 2180)	MEYDILTGYYGGYFDY (SEQ ID NO: 2179)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	SLATRPLGMDV (SEQ ID NO: 2184)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	DHFDTLTGYFRRLDS (SEQ ID NO: 2187)	VYYDILTGYNLFFDY (SEQ ID NO: 2177)	DAQSYYDILTGYQSYAFDI (SEQ ID NO: 2183)	VYYDILTGYNLFFDY (SEQ ID NO: 2177)	GPSTTYYDILTGYYTPYYYYYMDV (SEQ ID NO: 3014)	HVRDYDILTGYYRGHYFDY (SEQ ID NO: 2167)	ERGVVTAYGGDSFDL (SEQ ID NO: 2985)	DRGPGLLSSFFES (SEQ ID NO: 3033)	DEYYDILTGYQAPYYYYGMDV (SEQ ID NO: 3068)	ERGVVTAYGGDSFDL (SEQ ID NO: 2985)	DVTYHDIL TGYAGHEAFDI (SEQ ID NO: 3055)	ESGRYDILTGYYSGGGGMDV (SEQ ID NO: 3012)	DGANYDILTGYYTTTVYGMDV (SEQ ID NO: 3072)	RSYDILTGYYTYGMDV (SEQ ID NO: 3090)		KQRGDYDILTGYQLGYAFDI (SEQ ID NO	ERPGYDIL TGYPSSIYGMDV (SEQ ID NO: 3053)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	DTLGYDILTGYPPPYYYYDMDV (SEQ ID NO: 2988)	DTLGYDILTGYPPPYYYYDMDV (SEQ ID NO: 2988)	GRHYYDILTGYYNEAFDI (SEQ ID NO: 3031)	NYYDVLIQSYYGMDV (SEQ ID NO: 3077)
	99 - 120	99 - 108		•		99 - 115	99 - 112	99 - 114	99 - 114	99 - 114	99 - 109	98 - 113	98 - 113	99-113	99-113	99-117	99-113	99 - 123	100 - 118	99 - 113	99-111		99 - 113	98 - 116	102 - 121	99 - 119	99 - 114	99 - 107	100-119	99 - 118	99 - 114	99 - 120	95 - 116		99 - 113
51 - 66	20 - 66	20 - 66	20 - 66	20 - 66	49 - 65	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	49 - 62	49 - 65	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	52 - 67	20 - 66	20 - 66	20 - 66		20 - 65	52-69	20 - 66	20 - 66	20 - 66	25 - 67	20 - 66	20 - 66	20 - 66	46 - 62	50-65	20 - 66
26 - 36	26 - 35	26 - 35	26 - 35	26 - 35	26 - 34	26 - 35	26 - 35	26 - 35	26-35	26-35	26 - 35	26 - 34		26 - 35	26 - 35	26 - 35	26 - 35		26 - 37		26 - 35	•		26 - 35	26 - 37	26 - 35	26 - 35	26 - 35	26 - 37	26 - 35	26 - 35	26 - 35		26 - 35	26 - 35
1 - 128	1 - 131	1 - 119	1 - 124	1-117	1 - 124	1 - 126	1 - 123	1 - 125	1 - 125	1 - 125	1 - 120	1 - 124	1 - 124	1 - 124	1 - 124	1 - 128	1 - 124	1 - 134	1 - 129	1 - 124	1 - 122	1 - 130	1 - 124	1 - 127	1 - 132	1 - 130	1 - 125	1-118	1 - 130	1 - 129	1 - 125	1 - 131	1 - 127	. 1 - 126	1 - 124
1		223 - 231	230 - 239	223 - 233	231 - 241	233 - 243	230 - 240	231 - 240	231 - 240	227 - 237	226 - 235	231 - 241	230 - 240	229 - 240	230 - 239	230 - 240	230 - 239	236 - 246	234 - 244	229 - 239	228 - 237	•	•	233 - 242	236 - 244	236 - 245		224 - 233	235 - 245			•			230 - 239
- 201	- 204	1-190	٠.	84-190	91 - 197	94-200	91 - 197	92 - 198	92 - 198	88 - 194	87 - 193	92 - 198	- 197	: 961 - 06	91 - 197	91 - 197	191 - 197	97 - 203	95-201	90 - 196	- 195	- 202	- 197	•	97 - 203		•		196 - 202	96 - 202	•	ì	٠		191 - 197
62	28	- 168 18	175 19	- 168 18	- 175 19	- 178 19	- 175 19	- 176 19	176 19	- 172 18	- 171 18	- 176 19	- 175 19	- 174 19	- 175 19	- 175 19	-175 19	- 181 - 19	- 179	- 174 19	- 173	- 180	- 175 19	- 178 19	- 181	- 181	-176 1	- 169 1	- 180	- 180	-172 1	- 181	1771 -	-177	-175 1
166	169-	158	162	156	163	165	162	163	163	162	158	164	163	164	162	165	162	171	167	162	160	168	162	165	171	168	163	156	168	167	162		165	164	162
144 - 254	•	135 - 242	[40 - 250]	133 - 244	140 - 252	142 - 254	139 - 251	141 - 251	141 - 251	[41 - 248	136 - 246	140 - 252	140 - 251	140 - 251	140 - 250	144 - 251	140 - 250	150 - 257	145 - 255	140 - 250	138 - 248	146 - 256	140 - 250	143 - 253	148 - 255	146 - 256	141 - 251	134 - 244	146 - 256	145 - 255	141 - 248		143 - 253	1	140 - 250
-			_		•	_	1585 13			_	_	120051	•. •		_		•		•		_				1603	_	_			1 8091				1612 1	
_		_		1	-		7	_		-			-	_	_	_	-				,	_	~	_	_			~ 7		,-4		. •	. •	. ,	
I065D04	I065D08	1065F08	1067F05	1068B04	I068B08	1068C08	1068F03	1069B07	I071B03	I072B09	I073F04	I074B12	I075A02	1075G01	1078D02	1078D08	1078H08	I064A03	1064B03	1064B05	1064B11	1064C02	1064C03	1064C11	1064C12	1064D03	1064D04	1064D06	1064E05	1064E06	1064F07	1064F09	1064F10	1064F11	1064G01

(6)			
DNSGTYGY (SEQ ID NO: 3084) GGVTAGRSVYFDS (SEQ ID NO: 2990) SPNGDYSGYAWGLEY (SEQ ID NO: 3085) YFDGSGYYPVSFSY (SEQ ID NO: 3064) VYYDILTGLGYYFDY (SEQ ID NO: 3049) SYYDILTGRPYTDAFDI (SEQ ID NO: 2989) PLGITAVRGAKTDAFGI (SEQ ID NO: 2929) DRGASNYDILTGYYAPAQGVAFDI (SEQ ID NO: 2929)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) DGGGYDILTGYQYYYGMDV (SEQ ID NO: 2987) WATYYDTLTGYRLKDHAGFDI (SEQ ID NO: 3017) SPGDDILTGYYKYYFDY (SEQ ID NO: 3032) DAGESYDILTGYYVIEGYMDV (SEQ ID NO: 2986) EGAADYLNGOYFOH (SEO ID NO: 2985)	EGSWSGLDLDY (SEQ ID NO: 3007) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) VSGYNSGYFESYDMDV (SEQ ID NO: 2732) QGGQYDSPPLDV (SEQ ID NO: 3002) DRDYDILTDYSNYGMDV (SEQ ID NO: 3074) APLYDILTGYYYGGNDY (SEQ ID NO: 3028) DKDYDILTGYYYGGNDY (SEQ ID NO: 3040) DPNYDILTGYYYYAMDV (SEQ ID NO: 3062) EFDQLLARGHGMDV (SEQ ID NO: 3027) AGSSI MTYGTNV (SEQ ID NO: 3027)	ARGSYDILTGYYRPGDGYFDY (SEQ ID NO: 3043) ARGSYDILTGYYRPGDGYFDY (SEQ ID NO: 3043) GLYFEDTNYRHGDAFDI (SEQ ID NO: 2790) ERSYYDILTGYSPRSKYGMDV (SEQ ID NO: 3021) ATYDPLTGYSFDGFDI (SEQ ID NO: 2985) ERGVVTAYGGDSFDL (SEQ ID NO: 2985) RYSDALTGYSLGAFDV (SEQ ID NO: 2980) DYPIDVLTGYYPYGMDV (SEQ ID NO: 2860) DYPIDVLTGRRTKNWFDP (SEQ ID NO: 3013) DQVDRLLMQYNYYMDA (SEQ ID NO: 3047) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)
99 - 106 99 - 111 99 - 111 98 - 112 100 - 116 99 - 115	99 - 114 99 - 114 99 - 117 99 - 117 99 - 115 99 - 115	99 - 109 99 - 114 99 - 114 99 - 115 99 - 115 99 - 115 99 - 115	99 - 119 99 - 115 99 - 114 99 - 114 99 - 113 102 - 118 99 - 116
50 - 66 50 - 66 50 - 66 50 - 65 50 - 65 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 65 50 - 65 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
	26-35 26-35 26-35 26-35 26-35 26-35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35	26 - 35 26 - 35
1 - 122 1 - 122 1 - 123 1 - 123 1 - 124 1 - 126	1 - 125 1 - 125 1 - 125 1 - 128 1 - 129 1 - 126 1 - 130	1 - 125 1 - 125 1 - 125 1 - 121 1 - 126 1 - 126 1 - 126	1 - 130 1 - 130 1 - 125 1 - 125 1 - 124 1 - 129 1 - 127 1 - 125
	227 - 237 227 - 237 227 - 237 234 - 243 235 - 244 232 - 241 232 - 242	225 - 235 230 - 242 231 - 239 227 - 236 232 - 241 230 - 238 233 - 242 233 - 241 235 - 241	
189 195 200 200 202	20	192 193 193 193 193 193 193 193 193 193 193	
183 - 191 - 189 - 189 - 196 - 206 - 206 -	188 - 1 188 - 1 188 - 1 195 - 2 196 - 2 193 - 1 193 - 1	186 - 1 191 - 1 192 - 1 188 - 1 193 - 1 194 - 2 193 - 1	197 - 2 189 - 1 188 - 1 191 - 1 192 - 1 188 - 1
	162 - 172 162 - 172 162 - 172 166 - 179 167 - 180 164 - 177 161 - 174	158 - 170 163 - 175 163 - 175 159 - 172 164 - 177 165 - 178 165 - 178 160 - 170	
	248 16 248 16 248 16 254 16 255 16 253 16 249 16	246 15 253 16 225 16 247 15 252 16 249 16 252 16 252 16	
	141 - 2 141 - 2 141 - 2 144 - 2 145 - 2 146 - 2 139 - 2		
1614 1615 1616 1617 1618 1619 1620	1623 1623 1624 1625 1626 1628 1629	1630 1631 1632 1633 1634 1635 1636	1640 1641 1643 1644 1645 1646 1646 1649
1064G04 1064G08 1064G10 1064G11 1064H03 1064H06	1065A02 1065A04 1065A06 1065A07 1065B01 1065B09 1065B12	1065C02 1065C06 1065C08 1065C10 1065D01 1065D03 1065D06 1065E01	1065E06 1065E08 1065E09 1065E12 1065F04 1065F07 1065F09 1065F12

														_																					
								٠						3045					·			•													٠
30)				٠.		80)		<u>ଚ</u>	0:3009)			ন				97)			જ	_	(17	795)	3036)	360)		2736)			_		_		_ (6	
NO: 30	0:2983)	2984)	(05/	73)	: 2815)	NO: 30	0: 2153)	NO: 292	N CI OX	0: 2153)	348)	VO: 2992	(10	DV (SE	O: 3081)	NO: 29	0: 2153)	0: 2153)	NO: 305	0: 2153)	NO: 30	D NO: 2	ON A	O NO: 28			3054)	_	0: 2153)	_	0: 2153)	773)	0: 2985	NO: 292	773)
(SEQ ID	NOO	ON CHI	D NO: 7	NO: 27	OROLO	(SEO ID	ž A Ø	EQ ID	ÆV (SE	ž A O	NO.3	EQ ID	NO: 30	GGYYN	NO ID	(SEQ 11	Z A D S S	Z A OS	SEQ ID	Z O O	(SEQ ID	/ (SEQ I	S (SEQ	(SEQ 11	N D N	V (SEQ	ON CI	NO: 216	EO ED N	NO: 2	EQ 10 X	NO: 2	N A O	SECID	D NO: 2
GMDV	ADV (SI	'DI (SEC	F (SEQ	(SEQ II	OH (SE	AFDM	FDI (SI	AFGI (GRGEN	JFDI (SI	(SEQ 11	NFDY (S	(SEQ II	YYSAW	FDL (SI	RGMDV	SFDI (SI	GFDI (SI	GVDV (SFDI (SI	GMDV	(RYFD)	PSGYFI	YGMDV	FDP (SE		L (SEQ	SEQ ID	GFDI (SI	(SEQ II	GFDI (S)	(SEQ II	FDL (S	OAFGI (n das)
IGWVY	YYYD	YSGAF	RYYSD	YGTDV	NGQYF	LWERG	GYSFD	GAKTE	LTGYSI	GYSFD	IYGTDV	GYLSG	PPFDV	DILTG	IRGFGY	TGYYL	GYSFD(GYSFD(SYVYN	GYSFD(CTLGYF	LTGYS	LTGFY	TGYYP	TIGLGS	SPPYNY	RHAFD	GMDV (GYSFD	YGTDV	GYSFD	YGTDY	YGGDS	SGAKTI	YGIDV
DAYYDILTGWVYGMDV (SEQ ID NO: 3030)	FRYDILTGYYYDMDV (SEQ ID NO: 2983	EYYDILTGYSGAFDI (SEQ ID NO: 2984)	TRMDVLTRYYSDF (SEQ ID NO: 2750)	AGSSLMTYGTDV (SEQ ID NO: 2773)	EGAADYLNGQYFQH (SEQ ID NO: 2815	DTRVIGIQL WERGAFDM (SEQ ID NO: 3080)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153	PLGITAVRGAKTDAFGI (SEQ ID NO: 2929)	GRRYYDILTGYSLGRGEMDV (SEQ ID NO	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	AGTSLMNYGTDV (SEQ ID NO: 3048)	GPYDVLTGYLSGNFDY (SEQ ID NO: 2992)	QGGQYDSPPFDV (SEQ ID NO: 3001	GEKARYYDILTGYYSAWGGYYMDV (SEQ ID NO: 3045)	LNLEKTVIRGFGYFDL (SEQ ID NO: 3081)	VGGYDILTGYYLRGMDV (SEQ ID NO: 2997)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153	SPYDTLTGYVYNGVDV (SEQ ID NO: 3058	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	VAAAGARTLGYFGMDV (SEQ ID NO: 3071	DVSGHDILTGYSYRYFDV (SEQ ID NO: 2795	SPMYYDRLTGFYPSGYFDS (SEQ ID NO: 3036)	GAYYDIL TGYYPYGMDV (SEQ ID NO: 2860)	GPSSAGTTIGLGSFDP (SEQ ID NO: 3005)	ETRKYTSSPPYNYYYMDV (SEQ ID NO:	DOFSVGGRHAFDL (SEQ ID NO: 3054)	GMGDHYGMDV (SEQ ID NO: 2161)	ATYDPLTGYSFDGFDI (SEQ ID NO: 21	AGSSLMTYGTDV (SEQ ID NO: 2773)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2	AGSSLMTYGTDV (SEQ ID NO: 2773)	ERGVVTAYGGDSFDL (SEQ ID NO: 2985)	PLGITAVRGAKTDAFGI (SEQ ID NO: 2929)	AGSSLMTYGTDV (SEQ ID NO: 2773
	3 王		•	1	•	•		,		•	•		Ť.	-	$\overline{}$		•	•		4 A	14 V	_		_	•	_		_	_	•	•	110 A		115 P	∢ 0
101 - 116	99 - 11	99 - 113	99 - 111	99 - 110		99-115		99 - 115	98 - 117			99 - 114	_	_	99-11				99 - 114	99 - 11	99 - 11	99 - 11	99 - 117	_		99-11	94 - 10		99-11		99 - 11		$\overline{}$	7	99 - 11
50 - 68	51 - 66	50 - 66	50 - 66	20 - 66	20 - 66	20 - 66	99 - 09	99 - 05	50 - 65	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	99 - 09	20 - 66	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	45-61	20-66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66
26 - 35	26 - 36	26 - 35	26-35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 30	26-35	26 - 35	26 - 35	26 - 35	26-35	26-35	26 - 35	26 - 35
1 - 127	1 - 124	1 - 124	1 - 122	1-121	1 - 123	1 - 126	1 - 125	1 - 126	1 - 128	1 - 125	1 - 121	1 - 125	1 - 121	1 - 133	1 - 125	1 - 126	1 - 125	1 - 125	1 - 125	1 - 125	1 - 125	1 - 127	1 - 128	1 - 126	1 - 125	1 - 127	1-117	1-119	1 - 125	1 - 121	1 - 125	1 - 121	1 - 124	1 - 126	1 - 121
242	. 236	.236	.237	.236	.235	.241	.237	241	. 243	. 237	. 236	. 237	. 236	. 248	. 237	- 241	.237	. 237	. 240	.237	- 240	- 242	- 243	.241	237	- 239	- 232	-231	- 237	- 236	-237	- 236	- 239	- 242	- 236
233	226	226	228	227	225	232	227	232	233	227	227	227	227-	239	227	232	227	227	231.	227	230-	233	234	232	227	229 -	223	221	227	227	227	226	230	232	227
- 200	- 193	- 193	- 195	- 194	- 192	1- 199	- 194	- 199	1-200	1- 194	1-194	1-194	1	=	3 - 194	1-199	3 - 194	3 - 194	2 - 198	3 - 194	1-197	1-200	5 - 201	3 - 199		3 - 196	Ξ		3 - 194	8 - 194		7 - 193	1-197	3 - 199	8 - 194
		-	189	188	186	7 193	188	7 193	3 194	188	188	188	188	500	2 188	7 193	2 188	2 188	5 192	2 188	5 191	3 194	95	7 193	_		_	_	2 183	2 188		_	_	7 19.	2
- 178	•	171-	- 17	- 172	- 170	1-177	- 172	1-177	5-178	2-172	- 172	- 172	- 172	- 18	2-17	1-17	2-17	2 - 172	3 - 176	2 - 172	3 - 17.	5 - 178	5 - 179	1-17	_	1-174	$\overline{}$	5 - 166	2 - 17.	9-17	7	7	2-175	4 - 17	9 - 17.
165		161	Ξ.	159	160	164	162	26	166	162	159	_		171	3 162	164	3 162	3 162	163	3 162	163	3 165	166	_	_	164	_	2 156	8 162	7 159		_	_	_	7 159
3 - 253	140 - 247)-247	1	7-247	- 246	2 - 252	1-248	2-252	1-254	1-248	1-247	1-248	7-247	9 - 259	1-248	2 - 252	1-248	1 - 248	1-251	1 - 248	1-251	3 - 253	4-254	2 - 252	1-248	3-250	3 - 243	5 - 242	1 - 248	7-247	1 - 248	.•	0-250	1	7-247
143	14(140	138	137	139	142	141	142	4	141	137	141	137	149	141	142	141	141	141	141	141	143	144	142	141	143	133	135	141	137	141	137	140	142	137
1650	1651	1652	1653	1654	1655	1656	1657	1658	1659	1660	1661	1662	1663	1664	1665	1666	1667	1668	1669	1670	1671	1672	1673	1674	1675	1676	1677	1678	1679	1680	1681	1682	1683	1684	1685
6		5	. 4	Š.	وٍ		'n	_	9	0				æ	4	2	7	6	1	m	4	7	00		7	9	71	2	. 2	33	90	8	2	33	≱ .
1065G09	1065G10	1065H05	1065H07	1066A05	1066A06	1066A12	1066B05	1066B11	1066C06	1066C10	1066D02	1066D07	I066E01	1066E03	1066E04	1066E05	1066E07	1066E09	I066F01	1066F03	1066F04	1066F07	1066F08	1066F11	1066F12	1066G06	1066G07	1066H02	I067A02	I067A03	1067A06	1067A08	I067A10	1067B03	1067B04

																								6	•										
Diversity of the Cast of Cast	SCOOL MITVETTY (SEC. ID NO: 2962)	FOUND TOYOGEVEDY (SEC ID NO. 3011)	AGSSI MEVGEDY (SECTION: 3041)	TYYDII TGYSGGGAFDY (SEO ID NO: 2024)	GSRVRGVTPDI (SEO ID NO: 3024)	AGSSI MTYGTDV (SEO ID NO: 2772)	ECSGSSCPAROPPYYONYMIN (SEO ID NO: 2002)	AGSSI MTYGTDV (SEO ID NO: 2772)	GAYYDH TGYYDYGMDY (SBO ID NO: 2860)	OGGOYDSPPI DV (SEO ID NO: 3002)	AGSSLMTYGTDV (SFO ID NO: 2772)	GAYYDII.TGYYPYGMDV (SECTION NO. 2860)	DYRNYDILTGHPYYYGMDV (SEO ID NO. 2996)	OHYDILTGYSOEPFDI (SEO ID NO: 3022)	DOTYYDIL TGHYYYYGMDV (SEO ID NO: 3087)	EGAADYLNGOYFOH (SEO ID NO: 2815)	LGYYDILTGYRSDDY (SEO ID NO: 3029)	AGSSLMAYGTDV (SEO ID NO: 3016)	ENYDFLTGYYGAFDI (SEO ID NO: 2772)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	AGSSLMTYGTDV (SEO ID NO: 2773)	GGLYDILTGRPATDDAFDI (SEO ID NO: 3035)	TDRFGAKDVTARWGMDV (SEO ID NO: 2979)	GREDIDKVKPWDRYYHYYYMDV (SEQ ID NO: 2809)	DQGRYLDL (SEQ ID NO: 2175)	ELGLSIVGATTGALDM (SEQ ID NO: 2174)	ELGHREGGYWYSPYNV (SEQ ID NO: 2838)	KNMGASAADF (SEQ ID NO: 3042)	RYGDPFYYYYMNV (SEQ ID NO: 2755)	ESGSHYDLLTGLLVAANGFDV (SEQ ID NO: 3044)		MEYDIL TGYYGGYFDY (SEQ ID NO: 2179)	MEYDILTGYYGGYFDY (SEQ ID NO: 2179)	PYYDIL TGYFAFDI (SEQ ID NO: 3026)	MEYDILTGYYGGYFDY (SEQ ID NO: 2179)
90 - 106		99-114		99-115	99 - 109	99-110			99 - 115	99-110	99-110	99 - 115	99 - 117	99 - 114	99 - 117	99 - 112		99 - 110	99-113	99 - 114	99-110	99-117	99-115	99 - 120	99 - 106	98 - 113	99 - 114	98 - 108	99 - 112		99 - 114	99 - 114	99 - 114	•	99 - 114
50 - 66	50-66	20 - 66	20-66	50 - 66	50 - 66	50 - 66	50-66	20 - 66	20-66	50-66	50 - 66	50 - 66	50 - 66	50 - 66	20 - 66	99 - 05	50 - 66	50 - 66	99 - 09	20 - 66	20 - 66	99 - 05	99 - 09	20 - 66	20 - 66	49 - 65	20 - 66	20-66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66
26-35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35		26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 34	26-35	26 - 35	٠	26-35		26 - 35	26 - 35		26 - 35
1 - 117	1 - 121	1 - 125	1-121	1 - 126	1 - 120	1-121	1 - 130	1 - 121	1 - 126	1 - 121	1-121	1 - 126	1 - 128	1 - 125	1 - 128	1 - 123	1 - 124	1-121	1 - 124	1 - 125	1 - 121	1 - 128	1 - 126	1 - 131	1-117	1 - 124	1 - 125	1-119	1 - 123	1 - 130	1 - 125	1 - 125	1 - 125	1 - 123	1 - 125
224 - 233		231 - 240				223 - 233	235 - 245	223 - 233	232 - 241	227 - 236	227 - 236	231 - 241	234 - 243	229 - 237	234 - 243		226 - 236	227 - 236	226 - 237	•		230 - 240				1.	•			ŧ		•			227 - 237
- 191	- 194	- 198	- 194	- 199	- 193	184-190 2	202	190	- 199	- 194	- 194	- 198	201	961	201	192	193	194	193	198	193	- 197	- 200	- 205	- 193	- 198	- 198	161	961	204	96	194	196	961 -	- 194
59 185	_	76 192	72 188	77 193	11 187	168 184	180 196-	168 184 -	177 193	7 188	72 188	6 192	9 195-	4 190-	9 195	0 186-	1 187-	2 188-	1 187-	76 192-	1 187-					961 9			_		4 190-	2 188	4 190-	4 190	2 188
156 - 169	159 - 172	163 - 1	159 - 1	164 - 1	158 - 1'	158 - 10	168 - 18	158 - 10	164 - 17	159 - 17	159 - 17	64 - 17	166 - 17	64 - 17	166 - 17	160 - 17	161 - 17	159 - 17	161 - 17	7	59 - 17	7	7	•	57 - 169	71 - 501	7`	39 - 165	•	69 - 182	64 - 17	62 - 17	64-17	19 - 17	62 - 17
133 - 244	137 - 247	141 - 251	137 - 247	142 - 252	136 - 246	137 - 244	146 - 256	137 - 244	142 - 252	137 - 247	137 - 247	142 - 252	•	- 248	254	246	- 247	247	- 248	•	247	-251	-254	•	747	167-	727-	135 - 245	- 250	- 258	- 248	- 248	- 248	- 249	141 - 248
1686	1687	1688	1689	1690	1691	1692	1693	1694	1695	1696	1697	1698	1699	1700	1701	1702	1703	1704	1705	1706	1707	1708	1709	1710	1711	71/1	1713	1/14	C1/1	1/16	1717	1718	1719	07/1	17/1
I067C03	1067C05	1067C07	I067C10	I067C12	1067D01	1067D03	I067D05	1067D06	I067D09	I067D12	1067E02	I067E04	I067E05	1067F01	I067F03	I067F04	I067F08	1067F10	I067F11	1067G01	1067G09	1067H07	1068A07	1068E05	1008EU8 1069E13	1000011	1068504 106806	1069003	1000000	1008G11	1069A09	1069A10	1069B06	1009B09	7196001

	 MEYDILTGYYGGYFDY (SEQ ID NO: 2179) MEYDILTGYYGGYFDY (SEQ ID NO: 2179) GYYDILTGYYDAFDI (SEQ ID NO: 3051) DGYYDILTGYSGYYMDV (SEQ ID NO: 3059) DRLEYYDILTGYYYYYGMDV (SEQ ID NO: 3039) 	MEYDILTGYYGGYFDY (SEQ I MEYDILTGYYGGYFDY (SEQ I MEYDILTGYYGGYFDY (SEQ I SQSDYDILTGYYYYYYGMDV (S MEYDILTGYYGGYFDY (SEQ I MEYDILTGYYGGYFDY (SEQ I	MEYDLLTSYYGGYFDY (SQSDYDLLTGYYYYYYGN SQSNYDLLTGYYYYYYGN MEYDLLTGYYGGYFDY GMGDHYGMDV (SEQ ID	3 GMGDHYGMDV (SEQ ID NO: 2161) 5 AGTSLMNYGTDV (SEQ ID NO: 3048) 6 VPYYYDTSGGYLGEYYYGMDV (SEQ ID NO: 3010) 7 AGTSLMNYGTDV (SEQ ID NO: 3048) 7 ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) 7 SRDLLLFPHYGMDV (SEQ ID NO: 2133) 7 ATYDPLTGYSFDGFDI (SEQ ID NO: 2133)	0 1 1 1 0 1 1 0
	99 - 114 99 - 114 99 - 115 99 - 115	99 - 114 99 - 114 99 - 117 99 - 114	99 - 114 99 - 117 99 - 117 99 - 114 99 - 108	99 - 108 99 - 110 99 - 119 99 - 114 99 - 114	99 - 108 99 - 110 99 - 113 99 - 114 99 - 114 99 - 114
	50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 52 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35	26-35 26-35 26-35 26-35 26-35 26-35	26-35 26-35 26-35 26-35 26-35 26-35	26 - 35 26 - 35 26 - 37 26 - 35 26 - 35 26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35
1 - 127 1 - 127 1 - 126 1 - 126 1 - 124	1 - 125 1 - 125 1 - 124 1 - 126 1 - 129	1-125 1-125 1-128 1-128 1-125	1-125 1-128 1-128 1-125 1-119	1 - 119 1 - 121 1 - 130 1 - 121 1 - 123 1 - 123	1 - 119 1 - 121 1 - 125 1 - 124 1 - 119 1 - 125 1 - 125
1 1 1 1 1	227 - 237 227 - 237 226 - 236 228 - 238 231 - 241	227 - 237 229 - 237 227 - 237 234 - 243 231 - 240		225 - 234 227 - 236 236 - 245 227 - 236 231 - 240 229 - 238	225 - 234 227 - 236 231 - 240 228 - 238 224 - 234 227 - 237 231 - 240 224 - 234
196 195 193 193	2 2 2 2 8 2 2 8 8	194 198 198 198 198 198 198 198 198 198 198	198 197 188	192 194 198 198 198	192 198 195 191 194 198
	188 - 188 - 187 - 192 -	188- 190- 195- 192-		188 - 188 - 197 - 5 192 - 5 19	
164 - 174 164 - 174 163 - 173 163 - 173	162 - 172 162 - 173 161 - 173 163 - 173 166 - 176	162 - 172 164 - 174 162 - 172 166 - 179 163 - 176		157 - 170 159 - 172 168 - 181 159 - 172 163 - 176 161 - 174	157 - 170 159 - 172 163 - 176 162 - 173 157 - 169 162 - 172 163 - 176
1 1 1 1 1	- 248 - 248 - 247 - 249 - 252	- 248 - 248 - 248 - 254 - 251		247 - 247 - 247 - 247 - 251 - 249 - 251 -	-245 -247 -251 -249 -248 -251
143 142 143 140	14	14 14 14 14 14 14 14 14 14 14 14 14 14 1	14 4 4 4 5 E	135 137 146 137 139 139	135 137 141 140 135 141 141
1722 1723 1724 1725 1726	1727 1728 1729 1730 1731	1732 1733 1734 1735 1735	1738 1739 1740 1741 1742	1743 1744 1745 1746 1747 1748	1750 1751 1752 1753 1754 1755 1755
1069C06 1069C09 1069D03 1069E09	1069F05 1069F07 1069F12 1069G06 1069G08	1069G11 1070A03 1070A09 1070B01 1070B05	1070D04 1070D01 1070E01 1070G10 1071A06	1071B02 1071D02 1071D08 1071F01 1071G09 1072A01	1072B02 1072B10 1072B11 1072B12 1072C05 1072C10 1072D01

ATYDPL TGYSFDGFDI (SEO ID NO: 2153)	EGSYDILTGYYVGVGRMDV (SEO ID NO: 2171)	ATYDPL TGYSFDGFDI (SEO ID NO: 2153)	GMGDHYGMDV (SEO ID NO. 2161)	GMGDHYGMDV (SEO ID NO. 2161)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	DEVDILTGLIOGADY (SEO ID NO. 2883)	ATYDPLIGYSFDGFDI (SEO ID NO: 2153)	RDILTGEVDS (SEO ID NO: 2033)	GYRNDWYGAFFI (SEO ID NO: 3079)	ATYDPLIGYSFDGFDI (SEO ID NO. 2153)	ATYDPLIGYSFDGFDI (SEO ID NO: 2153)	AGTSLMNYGMDV (SEO ID NO: 3070)	GPYDILTGYYRDAFDI (SEO ID NO: 2998)	THYDILTGYYTADAFDI (SEO ID NO: 3019)	VOMDSEYYDLLTGINVGPYYFDY (SEO ID NO: 2132)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	GDFGDYDILTGYYPVYYGMDV (SEQ ID NO: 3082)	SYYDILTGYYPFGMDV (SEO ID NO: 3004)	DLWYYDILTGYYLDDAFDI (SEO ID NO: 2999)	DLWYYDILTGYYLDDAFDI (SEO ID NO: 2999)	SRDLLLFPHYGMDV (SEO ID NO: 2133)	TRMDVLTRYYSDF (SEQ ID NO: 2750)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	GYHDTLTSYNYNWFDP (SEQ ID NO: 3006)	AQMDSEYYDLLTGINVGPYYFDY (SEO ID NO: 3076)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	GMGDHYMDV (SEQ ID NO: 3008)	EMGYDILTGYYLNYMDV (SEO ID NO: 2862)	QHYDILTGYSQEPFDI (SEQ ID NO: 3022)	FNPTYDILTGYYIGGYFQH (SEQ ID NO: 2155)	ATYDPLTGYSFDGFDI (SEO ID NO: 2153)	ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)
99 - 114		99-114	99 - 108	99 - 108	99 - 114	99 - 113	99 - 114	101 - 110	99 - 110		99 - 114		99 - 114	99 - 115	99 - 121	99 - 114	99 - 114	99 - 114	99 - 119	99 - 114	99 - 117	99 - 117	99 - 112	99-111	99-114	99 - 114	99 - 114	99 - 121	99 - 114	99 - 107	99 - 115	99 - 114	101 - 119	99 - 114	99 - 114
50 - 66	20 - 66	50-66	20 - 66	99-09	99 - 09	99 - 09	50 - 66	50 - 68	20 - 66	50 - 66	50 - 66	99 - 09	99 - 09	99 - 09	50 - 66	20 - 66	99 - 09	20 - 66	50 - 66	20 - 66	99 - 09	99 - 0 9	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	99-05	20 - 66	20 - 66	20 - 68	20 - 66	20 - 66
26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35
1 - 125	1 - 128	1 - 125	1-119	1-119	1 - 125	1 - 124	1 - 125	1 - 121	1 - 121	1 - 125	1 - 125	1 - 121	1 - 125	1 - 126	1 - 132	1 - 125	1 - 125	1 - 125	1 - 130	1 - 125	1 - 128	1 - 128	1 - 123	1 - 122	1 - 125	1 - 125	1 - 125	1 - 132	1 - 125	1 - 118	1 - 126	1 - 125	1 - 130	1 - 125	1 - 125
231 - 240	234 - 243		221 - 231	221 - 231	231 - 240	226 - 236	227 - 237		226 - 236	231 - 240	231 - 240		229 - 237	232 - 241	238 - 247	231 - 240	231 - 240	231 - 240	232 - 242	229 - 237	234 - 243			•	•	•		234 - 244	32 - 241	•	232 - 241		•	231 - 240	231 - 240
192 - 198 2	201	- 198	- 188	188	198	193	- 194	- 193	- 193	198	198	194	961	199	202	198	198	86	66	96	201		192	195	86		194	201	199	187	661	194	203		- 198 2
	9 195	6 192	6 182	6 182-	6 192-	1 187-	2 188	1 187	1 187	6 192-	6 192-			_		6 192 -	6 192 -	6 192-]	7 193-1		_	_	_		_	_		_	_		_	Ξ.	1 197-	6 192	6 192
163 - 176	166 - 17	163 - 17	156 - 166	56 - 166	163 - 17	161 - 17	162 - 172	159 - 171	59 - 171	163 - 176	163 - 176	7	_	$\overline{}$	170 - 183	163 - 17	163 - 17	163 - 17	67 - 177	7	7	<u></u>	7	7	7	•	7	•	•	55 - 165	1	62 - 172	68 - 18	63 - 17	163 - 17
- 251	254	251	242	242 1	251 1	247 1	248 1	247 1	247 1	Ξ.	251 1	_	_	_		_	_	_	253 1		_	_	_	_	_	_	_	_	_	_	_	_	_	_	251 1
141-	144-	141.	135 -	135-	141 -	140 -	141 -	137-	137-	141 -	141 -	137-	141 -	142 -	148-	141 -	141 -	141 -	146 -	141 -	144 -	144 -	139 -	138.	•		141	148 -	141.	134-	•		•	٠	141
1758	1759	1760	1761	1762	1763	1764	1765	1766	1767	1768	1769	1770	1771	1772	1773	1774	1775	1776	1777	1778	1779	1780	1781	1782	1783	1784	1785	1786	1787	1788	1789	1790	1791	1792	1793
1072E01	I072E04	I072E05	1072E06	1072F03	1072F07	I072F11	I072G03	I072G04	I072G05	I072G09	I072H03	I072H07	I073A02	I073A03	I073A04	I073A05	I073A06	I073A09	I073A10	I073A11	I073B02	I073B05	I073B06	I073B07	I073B08	1073B11	1073C01	1073C02	1073C04	1073C07	1073C08	1073C09	1073C11	I073C12	1073D01

m -	QHYDLLTGYSQEPFDI (SEQ ID NO: 3022) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153) ENYDFLTGYYGAFDI (SEQ ID NO: 2772) ATYDPLTGYSFDGFDI (SEQ ID NO: 2153)	GEYDL.TGYPYWYFDL ATYDPL.TGYSFDGFDI ATYDPL.TGYSFDGFDI ATYDPL.TGYSFDGFDI DGSYDL.TGYYDNYM GEGGYDL.TGYL.GYL		ATYDPLIGYSFDGFDI (SEQ ID NO: 2133) ATYDPLIGYSFDGFDI (SEQ ID NO: 2153) ATYDPLIGYSFDGFDI (SEQ ID NO: 2153) ATYDPLIGYSFDGFDI (SEQ ID NO: 2153) TYYDILTGYYFDY (SEQ ID NO: 3056) ATYDPLIGYSFDGFDI (SEQ ID NO: 2153) ATYDPLIGYSFDGFDI (SEQ ID NO: 2153) LPPYDMLTGYYVGGGMDV (SEQ ID NO: 3050) AKPYTDFSRGSDADAFDV (SEQ ID NO: 3065)
	99 - 114 99 - 114 99 - 114 99 - 114 99 - 114	99 - 114 99 - 114 99 - 114 99 - 116 99 - 116		99-114 99-114 99-116 99-116 99-116
	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
26 - 35 26 - 35 26 - 35 26 - 35 26 - 37 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35		26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35
1-119 1-125 1-128 1-124 1-125 1-125 1-125	1 - 125 1 - 124 1 - 124 1 - 125 1 - 125 1 - 125	1 - 125 1 - 125 1 - 125 1 - 125 1 - 127 1 - 127	1-119 1-125 1-126 1-123 1-129 1-119	1 - 125 1 - 125 1 - 125 1 - 125 1 - 127 1 - 127
	231 - 240 231 - 240 230 - 239 230 - 240 231 - 240 231 - 240			221 - 240 227 - 237 231 - 240 231 - 240 224 - 234 231 - 240 234 - 244
	8 6 6 6 6 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	2002	192 192 193 193 193	192 - 198 2 188 - 194 2 192 - 198 2 192 - 198 2 195 - 201 2 195 - 201 3
-169 -176 -175 -175 -176 -176 -176 -176 -176 -176	- 176 - 176 - 175 - 175 - 175 - 176	175 176 176 175 178	- 169 - 172 - 173 - 170 - 170	172 172 176 176 169 176 177
2 = 4 0 = 8 = = =	-251 163 -251 163 -251 163 -251 163 -251 163 -251 163	-251 163 - -251 163 - -251 163 - -251 163 - -253 165 -		251 163 248 162 251 163 251 163 245 159 251 163 255 166 253 167
	141 140 141 141 141 141 141 141 141 141	141 141 141 141 143 14		141 141 138 141 143 143 143
1794 1795 1796 1797 1799 1800 1801	1803 1804 1804 1805 1806 1807	1809 1810 1811 1812 1813	1815 1816 1817 1818 1819 1820	1823 1824 1824 1825 1826 1827 1828
1073D03 1073D06 1073D08 1073D10 1073D11 1073E01 1073E02	1073E05 1073E06 1073E08 1073F01 1073F02 1073F03	1073F07 1073F11 1073F12 1073G03 1073G04	1073G05 1073G05 1073G07 1073G08 1073G09 1073G10	1073H01 1073H03 1073H05 1073H06 1073H07 1074A05 1074A06

		•	
DQGRYLDL (SEQ ID NO: 2175) RYGDPFYYYYMNV (SEQ ID NO: 2755) ELGLSIVGATTGALDM (SEQ ID NO: 2174) GGYDLTQYPAEFFHP (SEQ ID NO: 2174) DQGRYLDL (SEQ ID NO: 2175) DRYYDLTKGDYYYGMDV (SEQ ID NO: 3060) VQGETYYDLTGYWGPKRDLYGMDV (SEQ ID NO: 3069)	•	SPEGDYQPLSSNYNWLDP (SEQ ID NO: 3011) GKEGYNDN (SEQ ID NO: 3089) GSGYDLLTGYFTGSPLDY (SEQ ID NO: 2766) SPEGDYQPLSSNYNWLDP (SEQ ID NO: 3011) MGHYDILTGYFHYGMDV (SEQ ID NO: 2831) GNYDILTGYYHTPLDY (SEQ ID NO: 2831) SYYDILTGYYHTPLDY (SEQ ID NO: 3086)	GSGYDULIGYFIGSFLDY (SEQ ID NO: 2700) DDRDILTNYYLEYFQH (SEQ ID NO: 2868) GSGYDVLTGYFTGSPLDY (SEQ ID NO: 3057) GRYDILTGYFTSFDY (SEQ ID NO: 3066) DDRDILTNYYLEYFQH (SEQ ID NO: 2868) GTGYDILTGYYMGSAFDQ (SEQ ID NO: 2800) MGHYDILTGYYMGSAFDQ (SEQ ID NO: 2831) DQGRYLDL (SEQ ID NO: 2174) GTGYDILTGYYMGSAFDQ (SEQ ID NO: 2174) GTGYDILTGYYMGSAFDQ (SEQ ID NO: 2800) DQGRYLDL (SEQ ID NO: 2175) RDVQGAPY (SEQ ID NO: 2175)
99 - 106 99 - 112 98 - 113 99 - 114 99 - 106 99 - 123	98 - 113 99 - 106 98 - 113 101 - 119 99 - 110 99 - 106	99 - 116 99 - 116 99 - 116 99 - 115 99 - 111	99-116 99-116 99-117 99-116 99-116 99-106 99-106
50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	49 - 65 50 - 66 50 - 66 49 - 65 50 - 68 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 51 - 66 51 - 66
26-35 26-35 26-35 26-35 26-35 26-35	26 - 34 26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35	26 - 35 26 - 3
1 - 117 1 - 123 1 - 124 1 - 125 1 - 117 1 - 127	1 - 124 1 - 122 1 - 124 1 - 120 1 - 121 1 - 121 1 - 126	1-127 1-127 1-127 1-126 1-126	1-125 1-125 1-127 1-127 1-127 1-127 1-127 1-127
221 - 231 230 - 240 230 - 240 230 - 240 224 - 235 232 - 242 241 - 251	230 - 240 229 - 238 224 - 234 230 - 240 237 - 247 225 - 233 233 - 243 223 - 231		232 - 242 233 - 243 233 - 243 232 - 243 232 - 242 233 - 242 231 - 241 234 - 244 222 - 233
- 188 - 197 - 197 - 191 - 199 - 208	191 - 197 185 - 191 191 - 197 191 - 197 198 - 204 186 - 192 194 - 200 184 - 190	200 199 198 198	193 - 199 193 - 199 194 - 200 190 - 196 193 - 199 193 - 199 195 - 201 185 - 191
-166 -175 -175 -175 -169 -177	-175 -174 -169 -175 -170 -170 -178	177 177 174 175 176 177 177 176 177 176	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
242 156 251 162 251 163 251 163 251 165 246 156 253 167 262 173	251 163 - 249 161 - 245 156 - 251 163 - 258 169 - 244 160 - 254 166 - 242 158 - 244 157 - 244 157 - 244 157 - 244 157 - 249		253 107 254 164 255 166 253 167 253 167 255 165 255 163 243 157 245 156
133 - 140 - 140 - 141 - 141 - 150 -	140 - 138 - 138 - 140 -		143 - 143 -
1830 1831 1832 1833 1834 1834 1835	1837 1838 1839 1840 1841 1842 1843 1844	1846 1847 1848 1849 1850 1851	1854 1855 1856 1857 1858 1859 1860 1861 1862 1863 1863
1074B03 1074B11 1074C07 1074D03 1074D04 1074D07	1074D08 1074D11 1074E05 1074E07 1074E09 1074E11 1074H05 1075A03	1075B07 1075D11 1075D12 1075G02 1075G09 1075G10	10/5H0/ 1076A11 1076A12 1076B10 1076B12 1076C06 1076C01 1076E08 1076E08

																																	•		
	_		ন																											٠.			•		
VEGVYDILTGYSFDAFDI (SEQ ID NO: 3078)	EQGYDILTGYYPEGGWFDP (SEQ ID NO: 2834)		•	MEYDILTGYYGGYFDY (SEQ ID NO: 2179)	EKYDILTGYYDAFDI (SEQ ID NO: 3046)	EMGYDILTGYYLNYMDV (SEQ ID NO: 2862)	EMGYDILTGYYLNYMDV (SEQ ID NO: 2862)	MEYDILTGYYGGYFDY	MEYDILTGYYGGYFDY (SEQ		MEYDILTGYYGGYFDY (SEQ ID NO:	ESHYDILTGYYSNPSFDI (SEQ		TGSGFDY (SEQ ID NO: 2192)	Ξ.	DWDMDV (SEQ ID NO: 2193)		_	•		_	•		VHSTGYAFEN (SEQ ID NO: 2200)	_	VGIKAAAVDNFEY (SEQ ID NO: 2197)	_					DASKDIVVLPLAI (SEQ ID NO: 2198)			
99 - 116		98 - 113	102 - 120		99 - 114	99 - 114	99 - 114	99 - 113	99 - 115	99 - 115	99 - 114					99 - 110	99 - 105	99 - 109	99 - 104	99 - 109	99 - 103	99 - 107	99 - 104	99 - 107	99-111	99 - 111	99 - 108	99 - 115	. 99-111	99 - 114	99 - 109	99 - 109	99 - 104	99-110	99-111
50 - 66	20 - 66	49 - 65	52 - 69	20 - 66	20 - 66	20 - 66	99 - 09	99 - 09	20 - 66	99 - 05	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	99 - 05	20 - 66	20 - 66	20 - 66	20 - 66	99-05	20 - 66	99 - 09	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66			20 - 66	20 - 66	20 - 66	20-66	20 - 60
26 - 35	26 - 35	26 - 34	26 - 37	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35
1 - 127	1 - 128	1 - 124	1-131	1 - 125	1 - 125	1 - 125	1 - 125	1 - 124	1 - 126	1 - 126	1 - 125	1 - 125	1 - 125	1 - 125	1 - 127	1 - 121	1 - 116	1 - 120	1-115	1 - 120	1-114	1 - 118	1-115	1-118	1 - 122	1 - 122	1 - 119	1 - 126	1 - 122	1 - 125	1 - 120	1 - 120			1 - 122
233 - 243	233 - 243	229 - 239	237 - 246	231 - 240	231 - 240	227 - 237	231 - 240	226 - 236	232 - 241	232 - 241	229 - 237	227 - 237	231 - 240	229 - 237	233 - 242	225 - 233	220 - 228	224 - 232	222 - 231	227 - 236	220 - 229	222 - 230	221 - 230	222 - 230	227 - 237	228 - 238	226 - 236		228 - 238	232 - 242	•		1	•	226 - 234
194 - 200	94 - 200		98 - 204	192 - 198	192 - 198	188 - 194	192 - 198	187 - 193	193 - 199	193 - 199	190 - 196	188 - 194	192 - 198	190 - 196	94 - 200	186 - 192	181 - 187	185 - 191	183 - 189	88 - 194	81 - 187	83 - 189	182 - 188	183 - 189	88 - 194	89 - 195	187 - 193	194 - 200	189 - 195	193 - 199	188 - 194		181 - 187	187 - 193	187 - 193
6-178	8-178	-174	- 182	- 176	-176	-172	- 176	- 171	- 177	- 177	- 174	-172	- 176	-174	-178 1	- 170	- 165	- 169	- 167	- 172	- 165 1	- 167	- 166	-167	160 - 172 1	- 173 1	_	_	- 173	164 - 177 1	- 172	- 172	- 165	171 -	1-171
4 16	4 16	0 164	7 169	1 163	1 163	8 162	1 163	7 161	2 164		8 164	8 162	1 163	8 164	3 165	4 160	9 155	3 159	2 154	7 159	0 152	1 157	1 153	1 157	_	19 161	_	_				_	_		191
143 - 254	144 - 254	140 - 250	147 - 257	141 - 25]	141 - 251	141 - 248	141 - 25	140 - 247	142 - 252	142 - 252	141 - 248	141 - 248	141 - 251	141 - 248	143 - 253	137 - 244	132 - 239	136 - 243	131 - 242	136 - 247	130 - 240	134 - 241	131 - 241	134 - 241	138 - 248	138 - 249	135 - 247	142 - 254	138 - 249	141 - 253	- 1		•	• 1	138 - 24:
1865	1866	1867	1868	1869	1870	1871	1872	1873	1874	1875	1876	1877	1878	1879	1880	1881	1882	1883	1884	1885	1886	1887	1888	1889	1890	1891	1892	1893	1894	1895	1896	1897	1898	1899	1900
1076G01	1076H01	1076H03	I077B05	I077C10	1077D01	I077D04	1077D11	1077D12	I077E01	I077E03	I077E08	I077F05	1077G06	I077H02	I078B05	1079E02	I079F11	1082G02	1082H08	I099D03	1079B05	I079B12	1079C01	I079F06	1079F08	I080A03	1080A08	I080B01	1080D03	1080E05	1080G07	1080G09	I082A05	I082B08	I082C03

		·	
- 107 WTSSGAFDI (SEQ ID NO: 2205) - 111 DRGSGWPNWYFDL (SEQ ID NO: 2212) - 110 ESGAGGYYYDDY (SEQ ID NO: 2196) - 111 VGIKAAAVDNFEY (SEQ ID NO: 2197) - 103 DTTDY (SEQ ID NO: 2203) - 104 DTTDY (SEQ ID NO: 2203) - 104 NLWGLDY (SEQ ID NO: 2211) - 107 GNAWGAFDI (SEQ ID NO: 3123) - 107 GGMDWDFDY (SEQ ID NO: 3123) - 107 VDSSGYAYY (SEQ ID NO: 3123) - 107 VDSSGYAYY (SEQ ID NO: 3213)	121 109 107 113	112 113 110 110 110 108	 110 EAYTSSWAEFDF (SEQ ID NO: 3190) 109 NITPLAMVGDF (SEQ ID NO: 3146) 103 LEDF (SEQ ID NO: 3161) 104 DSGSPD (SEQ ID NO: 3108) 107 EGVAAGEDY (SEQ ID NO: 3123) 108 EKRGSRRVFDI (SEQ ID NO: 3193) 110 EAYASSWAEFDF (SEQ ID NO: 3189) 110 PYGSGSYAFDI (SEQ ID NO: 3185) 110 ARDYYDSSGYYVPDAFDI (SEQ ID NO: 3107)
99 - 107 99 - 111 99 - 1110 99 - 1111 99 - 103 99 - 107 99 - 107 99 - 107	99 - 109 101 - 121 99 - 108 99 - 107 99 - 109 99 - 103	99 - 112 99 - 117 99 - 113 99 - 110 99 - 110 99 - 108	99 - 110 99 - 109 99 - 103 99 - 104 99 - 107 99 - 109 99 - 109
26 - 35		****	50 - 50 - 50 - 50 - 50 - 50 - 50 - 50 -
26.25.25.25.25.25.25.25.25.25.25.25.25.25.	26-35 26-35 26-35 26-35 26-35 26-35	26-35 26-35 26-35 26-35 26-35 26-35 26-35	26 - 35 26 - 35
1-118 1-122 1-122 1-122 1-114 1-115 1-118 1-118 1-118	1 - 120 1 - 132 1 - 119 1 - 118 1 - 120 1 - 114	1 - 123 1 - 128 1 - 116 1 - 124 1 - 121 1 - 128 1 - 119	1 - 121 1 - 120 1 - 114 1 - 118 1 - 118 1 - 120 1 - 121 1 - 121
222 - 230 226 - 234 226 - 236 228 - 238 219 - 229 218 - 226 217 - 227 224 - 233 220 - 230	225 - 235 234 - 244 223 - 231 223 - 232 225 - 235 220 - 229		226 - 236 222 - 232 220 - 239 217 - 227 222 - 230 223 - 236 224 - 236 232 - 242
- 189 - 193 - 193 - 195 - 195 - 184 - 191 - 191 - 191			- 183 - 184 - 184 - 190 - 193 - 191
			183 183 178 178 183 184 187 187
157 - 167 161 - 171 161 - 171 161 - 173 152 - 164 153 - 163 156 - 169 156 - 169 156 - 169 156 - 169 156 - 169 156 - 169	158 - 170 169 - 179 158 - 168 156 - 168 158 - 170 152 - 165	162 - 172 167 - 177 155 - 163 163 - 173 158 - 168 166 - 179 157 - 169	159 - 171 157 - 167 152 - 165 152 - 165 157 - 167 158 - 168 159 - 169
241 243 244 244 244 241	246 242 243 243 246 246	246 251 239 244 254 245 243	243 243 241 244 247 253
_	136 - 148 - 135 - 136 - 130 -		136 - 136 - 136 - 136 - 136 - 136 - 136 - 136 - 136 - 143 -
1901 1902 1903 1904 1906 1907 1908 1910 1911	1913 1914 1915 1916 1917 1918	1920 1921 1922 1924 1926 1926	1928 1930 1931 1932 1933 1934 1935
1082D07 1082G01 1083B12 1083B03 1084A01 1084B02 1084C04 1079A01 1079A03 1079A04	1079A07 1079A10 1079A11 1079B02 1079B03 1079B04	1079B09 1079C02 1079C04 1079C05 1079C07 1079D01 1079D04	1079503 1079503 1079508 1079501 1079506 1079511

			٠.																																
		(612																																	
		LPPDLKYCDGGMCSGFDWLGP (SEQ ID NO: 3219)	િ											í	100)	(8)	3	3140)				<u>د</u>		-	<u>-</u>				DYYDGSSYSSGDYYYYWWV (SFO ID NO. 3227)	3113)	(6116	•			
		P (SEQ I	ESLLITEEYCGSDCYS (SEQ ID NO: 3115)	(660		146)	3166)	146)	124)	(8)	192)	_			ESECUTATION VALITATION (SEQUED NO; 3160)	PARTICIONAL PROPERTY (SEQ ID NO: 3188)	(c)	·· /	3181)	: 3096) :	3200)	COVOLCI I I I I IMD (SEQ ID NO: 3125)	á	RKATETYSGEANDEDY (SECTION)	100.015 100.0	(2016	2:3109)	0:3001)	FO ID	DSDLVVIPTAIOGRYYFDN (SEO ID NO: 3113)	3130	(222	3155)	: 3153)	(220)
	GHFYGMDV (SEQ ID NO: 3098)	FDWLG	SEQ ID	NSAPPAPSMDV (SEQ ID NO: 3099)	139)	NITELAM GDF (SEQ ID NO: 3146)	AD I SIND I YMDV (SEQ ID NO: 3166)	NITELAM VGDF (SEQ ID NO: 3146)	FLESY Y YMDV (SEQ ID NO: 3124)	GNSFGKILDY (SEQ ID NO: 3158)	DVFFFDGYLEV (SEQ ID NO: 3192)	(C: 31/1	3210)	7: 3111)	7 (SEC.)	FACUSOS I AFFERT (SEQ ID N	15 :5N V	CONTRACTOR OF THE PROPERTY (SECTION NO. 2162)	MAGGVEST GEDN (SEQ ID NO: 3181)		AVPSBGVVVVVVAAADV (SEQ ID NO: 3206)		7: 3141)	REALTERSON (SEC ID NOT 3100)	RPAT RSI WATER (SEC ID NO:	HCTGGSCGF (SEO ID NO: 3185)	NPYYYDSSEGFFDY (SEO ID NO: 3109)	SGROAYYYYGMDV (SEO ID NO: 3091	S) ACIVAL	DN (SEC	GKRYSYGWYFDI (SEO ID NO: 3130)	(46)	EGDPTDNDAFDV (SEQ ID NO: 3155	DGPTYARPYYLDH (SEO ID NO: 3153	DGTKYDWGFDY (SEQ ID NO: 3220)
	SEQ ID 1	GMCSG	SDCYS (V (SEQ	KYYDY (SEQ ID NO: 3139)	r (SEQ.	OHS) AC	F (SEQ 1	V (SEQ		V (SEQ.)		BUSSEDV (SEQ ID NO: 3210)	ERSSSFD1 (SEQ ID NO: 3111) BGBGDGVVNA PROVIENS	ער ז ז רט פרטונים	SEFECT Y	77(2)	TWO YOU		שכי) ארוז ספיטיי זכו	730,77		WANTELLI (SEQ ID NO: 3141)	ייושפרוי הפהרוי	(650)	CEO TO	FEDY (S)	MDV (S	WYYOU	CRYY	DI (SEO	DTPLDP (SEQ ID NO: 3094)	ON (SEO	LDH (SE	Y (SÈQ
	GMDV(KYCDC	EEYCG	AFSMU	Y (SEQ.	AIM V GL	MY YM	AMVGU	XXXMI	ikilluy Tagara	DOYLE BEDY (יייייייייייייייייייייייייייייייייייייי	Tac) and	ייוקעקע	ANT DO	VVEDV			C LLOON	777777		11110 100 VCI	ir Di Vytehyv	TYSGE/	OI WAR	TONO!	DSSEG	YYYYG	3SYSS(VIPTAI	YGWYF	P (SEO I	DNDAFI	ARPYY	DWGFE
•	-				KYYD	MILL	AUrs	NIIPL	rrres Sign	COSFC	ACVO	TICK TATO	DELEGG	ECCTO TOTOT				I GPAN		1470	AVOCA	Idadas	Tagos.	RKATE	RPATR	1.HCTG	NPYYY	SGROA	DYYDO	DSDLV	GKRYS	DTPLD	EGDPT	DGPTY	DGTKY
	99-100	171 - 171	99 - 113	201 - 20	201 - 20	100	29 - 109			207 - 60	99 - 107	90-104	90-104	99-114	00 - 114	90-108	90-114	99-111	90-111	00-110	107 - 115	90-106	100 - 100	99 - 115	100-111	101 - 110	99 - 112	99-111	99-117	99 - 117	99-110	99 - 104	99-110	99-111	99 - 109
77 03	20 - 00	00-00	00 - 00	00-00	99-09	20-02	20-05	90-00	99-05	20-05	20 - 66 50 - 66	20-66	50.66	50 - 66	50 - 66	20 - 66	20 - 65	20 - 66	20 - 66	20 - 66	52 - 69	20- 50-	52-67	99-09	52 - 67	20 - 68	99 - 09	20 - 66	51 - 66	99-05	99 - 0 5	20 - 66	99-09		99 - 09
36 36		26 26	26 - 35	26 - 26	26-35	36.36	26 26	26.35	26 - 25	26 - 25	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35		35	35	37	35	3.6	35	37	26-35	26 - 35	35	26 - 36	26-35	26 - 35	35	35	35	26-35
1-117	1 - 117	1 1 1 2 4	1 - 124	1 - 120	1 - 120	1 - 120	1, 120		1-110	1.120	1-118				1 - 125	1-119															- 121 -				- 120
219-230		256-256	220 - 230			٠,	٠.					1	•	2 - 242	•		1 - 241					2 - 232	•		228 - 238	227 - 237		•		•	3-238		3 - 238	3-238	5-234 1
- 186 21				•	•				2	189	189	188	191	199	198	193	861	7	195	195 228				200	195 22	194 22	201	194	8	• •	• •		• • •	•	- 193 226
180	_	187	183	12	183	185	183	185	186-	183	183 -	182-	185-	193-	192	187	192 - 198	188 -]	189 - 195	189 - 195	193 - 199	183 - 189	188-	194	189-	188 -	195	188 -			189	183 -	- 681	68 5	187 -
154 - 164	169 - 179		•		157 - 167	- 1		•			57 - 167	54 - 166	156 - 169	64 - 177	921 - 99	158 - 171	164 - 176	162 - 172	161 - 173	161 - 173	165 - 177	57 - 167	59 - 172	165 - 178	161 - 173		64 - 179	ı	•	67 - 180	7	154 - 167	160 - 173	•	101 - 171
- 241	253	247	243	237	243	243	243	243		_	241 1	242 1	245 1	_	250 1	247 1	252 1	248 1	249 1	249 1	254 1	243 1	248 1	254 1	49	_		_ '		_ '		_ `			245
133 -	148-	140 -	136-	130 -	136 -	136-	136-	136-	135 -	136-	•		133 -		•		•	•	138	•	•	•	136-2	•	138-	•	1				•	•	•	138-2	
1937	1938	1939	1940	1941	1942	1943	1944	1945	1946	1947	1948	1949	1950	1951	1952	1953	1954	1955	1956	1957	1958	1959	1960	1961	1962	1963	1964	1900	1700	1961	1908	1969	1970	17/1	7/61
1079F01	1079F02	I079F03	1079F04	I079F09	I079F10	I079F12	I079G02	1079G05	1079G06	I079H05	1079H06	\A01	A02	A05	1080A06	A07	A10	B02	B03	B05	B06	B07	B08	B09	B10	B11	812 703	ָה מ) å	9 8	3 5	3 5	7 6	3
107	107	1075	107	I075	· 1075	107S	1075	1079	107s	1079	1079	I080A01	I080A02	I080A05	1080	I080A07	I080A10	I080B02	1080B03	1080B05	1080B06	I080B07	I080B08	1080B09	1080B10	I080B11	1080B12			1030C0 1080C0	000	108001	100001	100001	7007

•		6	
HETFSHCSGGSCYPFDY (SEQ ID NO: 3212) SGRQAYYYYGMDV (SEQ ID NO: 3091) HETGYVYLTDY (SEQ ID NO: 3165) LHCTGGSCGF (SEQ ID NO: 3186) VDYTDYEMGAFEI (SEQ ID NO: 3187) VDYTDYEMGAFEI (SEQ ID NO: 3187) SSRNGGDY (SEQ ID NO: 3214)			111 ERGGRDGDYALDF (SEQ ID NO: 3148) 112 RTPDHNGDSGPPDY (SEQ ID NO: 3215) 103 DTTDY (SEQ ID NO: 2203) 108 ESLTGGAFDI (SEQ ID NO: 3117) 103 DTTDY (SEQ ID NO: 2203) 103 DTTDY (SEQ ID NO: 2203) 104 GAGSRYFDL (SEQ ID NO: 2203)
99 - 114 99 - 111 99 - 111 99 - 110 99 - 110 99 - 111 99 - 108	99 - 109 100 - 109 101 - 110 99 - 115 99 - 103 99 - 111 99 - 113 99 - 103	99 - 116 100 - 109 100 - 113 99 - 112 99 - 110 100 - 109 99 - 110	99 - 111 99 - 112 99 - 103 99 - 103 99 - 103 99 - 103
666	50 - 66 52 - 67 50 - 68 50 - 66 50 - 66 50 - 66 51 - 66 52 - 66	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66	50 - 66 51 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
26 - 35 26 - 35 26 - 35 26 - 35 26 - 35 26 - 35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35	26-35 26-37 26-37 26-35 26-35 26-35 26-35 26-35	26-35 26-35 26-35 26-35 26-35 26-35 26-35 26-35
- 125 - 120 - 120 - 121 - 122 - 119	- 120 - 121 - 121 - 121 - 122 - 124 - 124	- 127 - 127 - 124 - 124 - 123 - 119 - 121 - 121	1-122 1-123 1-114 1-119 1-114 1-114 1-118
199 232 - 194 227 - 192 225 - 196 229 - 193 226 -	196 196 196 196 196 196	199 232 - 199 232 - 199 232 - 199 232 - 198 231 - 192 227 - 192 228 - 195 228 - 195 228 - 195 228 - 194 227 - 197 22	- 197 230 - - 195 228 - - 185 218 - - 192 225 - - 185 218 - - 183 216 - - 186 219 -
193 188 186 188 190 187	186 187 187 187 188 189 180 180	193 187 192 188 188 186 189 189 189	191 189 179 179 177 180 181
	160 - 170 159 - 171 160 - 172 165 - 178 160 - 173 160 - 173 161 - 173 162 - 177 164 - 164		
	136 - 246 136 - 247 137 - 248 142 - 254 130 - 242 138 + 249 142 - 253 138 - 248 140 - 250		
1973 1974 1975 1976 1977 1978	1980 1981 1982 1983 1984 1986 1987 1989 1989	1992 1993 1994 1995 1996 1997 1999	2001 2002 2003 2004 2005 2006 2007
1080D02 1080D04 1080D05 1080D08 1080D09 1080D11	1080E01 1080E04 1080E04 1080E07 1080E07 1080F04 1080F05 1080F06	1080G04 1080G10 1080G11 1080H01 1080H03 1080H04 1080H05 1080H06	1080H08 1080H09 1081A01 1081A04 1081A06 1081A08

																	•																	
	ı																						D: 3184)											-
GGDRAFDI (SEQ ID NO: 3119)	GNAWGAFDI (SEQ ID NO: 2211)	GGDRAFDI (SEQ ID NO: 3119)	VKRYYFDY (SEQ ID NO: 3179)	ELTGANDAFDI (SEQ ID NO: 3104)	RRYALDY (SEQ ID NO: 2920)	DTTDY (SEQ ID NO: 2203)	DTTDY (SEQ ID NO: 2203)	GFALYKD (SEQ ID NO: 3169)	DTTDY (SEQ ID NO: 2203)	DTTDY (SEQ ID NO: 2203)	EDLTGDAFDI (SEQ ID NO: 3103)	GDAYFDY (SEQ ID NO: 3147)	GDAYFDY (SEQ ID NO: 3147)	DTTDY (SEQ ID NO: 2203)	DTTDY (SEQ ID NO: 2203)	EGLLDAFDI (SEQ ID NO: 3200)	DTTDY (SEQ ID NO: 2203)	VGYGGKGDY (SEQ ID NO: 3137)	GAGSRYFDL (SEQ ID NO: 3118)	GLAPIVDGGMTNDAFDI (SEQ ID NO: 3184)	DTTDY (SEQ ID NO: 2203)	RLIRKAR (SEQ ID NO: 3170)	DTTDY (SEQ ID NO: 2203)	ERGNQAFDI (SEQ ID NO: 3156)	RRYALDY (SEQ ID NO: 2920)	DTTDY (SEQ ID NO: 2203)	DTTDY (SEQ ID NO: 2203)		SRSPYDAFDI (SEQ ID NO: 3097)	(SEQ ID NO:	DTTDY (SEQ ID NO: 2203)			
99 - 106		99 - 106		99 - 109	99 - 105	99 - 103	99 - 103	99 - 105	99 - 103	99 - 103	99 - 108	99 - 105	99 - 105	99 - 103	99 - 103	99 - 107	99 - 103	99 - 103	99 - 103	99 - 103	99 - 107	99 - 107		99 - 103	99 - 105		99 - 107				•		•	99 - 103
50 - 66	99 - 09	20 - 66	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	99 - 09	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	99 - 09	20 - 66	99-05	99 - 09	99 - 09	20 - 66	20 - 66	20 - 66		20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66	20 - 66
26-35		26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35				26 - 35		26 - 35	26 - 35		26 - 35	ı			•	•	26 - 35	26 - 35
1-117	1-118	1 - 117	1 - 117	1 - 120	1-116	1 - 114	1-114	1 - 116	1-114	1 - 114	1-119	1-116	1 - 116	1 - 114	1 - 114	1 - 118	1 - 114	1 - 114	1 - 114	1 - 114	1 - 118	1 - 118	1 - 126	1 - 114	1-116	1 - 114	1 - 118	1 - 116	1 - 114	1-114	1 - 114	1 - 119	1-114	1 - 114
223 - 232		223 - 232	219 - 229	222 - 232	218 - 228	219 - 229	218 - 226	218 - 228	218 - 226	218 - 226	221 - 231	218 - 228	218 - 228	217 - 227	219 - 229	224 - 233	218 - 226	218 - 226			220 - 230	220 - 230		219 - 228		1	i			•	•	•	218 - 226	219 - 229
184 - 190	1	184 - 190	180 - 186	183 - 189	179 - 185	180 - 186	179-185	179 - 185	179 - 185	179 - 185		•	•		180 - 186	185 - 191		179 185		•	•	181 - 187		180 - 186			1	•					•	180 - 186
155 - 168	•	155 - 168		157 - 167	153 - 163	152 - 164	153 - 163	153 - 163	153 - 163	153 - 163	•	7	153 - 163	152 - 162	7	156 - 169	•	153 - 163	152 - 164		155 - 165	155 - 165	163 - 173	152 - 164	153 - 163	•		153 - 163		4	1	157 - 170	153 - 163	152 - 164
133 - 243 130 - 236	134 - 244	133 - 243	133 - 240	•	132 - 239	130 - 240	130 - 237	132 - 239	130 - 237	130 - 237	135 - 242	132 - 239	132 - 239	130 - 238	130 - 240	134 - 244	130 - 237	130 - 237			•	134 - 241	142 - 249		-		•		•			135 - 245		130 - 240
2009	2011	2012	2013	2014	2015														2029	2030	2031	2032	2033	2034	2035	2036	2037	2038	2039	2040	2041	2042	2043	2044
1081A10 1081B01	I081B04	I081B05	I081B06	I081B07	I081B08	I081B09	I081B10	I081B11	1081C07	I081C08	1081D04	1081D06	I081D08	I081D09	1081D10	1081D11	I081D12	1081E02	1081E03	1081E05	1081E06	1081E07	1081E10	1081F01	1081F04	I081F05	1081F06	I081F07	I081F11	1081G01	1081G04	1081G06	1081G10	1081H02

															•						ପ	•				. •									133)
): 2203)	O ID NO: 3202)	:3174)): 2203)	PAASSRGPKDAFDI (SEO ID NO: 3129)	: 3122)	JD NO: 3123)	2210)	NO: 3135)	1D NO: 3123)		SEO ID NO: 3143)	(O ID NO: 3093)	SEO ID NO: 3106)	: 2210)	(ID NO: 3152)	3167)	VEWEDIVVGSAFDI (SEQ ID NO: 3128)	EO ID NO: 3177)	SEO ID NO: 3166)	ID NO: 3123)	GPIYYFDGSAYEGYYFDY (SEQ ID NO: 3222)) NO: 3223)	I (SEQ ID NO: 3129)	3Q ID NO: 3110)	() ID NO: 3186)	Q ID NO: 3173)	(SEQ ID NO: 3138)	VDYTDYEMGAFDL (SEQ ID NO: 3172)	(SEQ ID NO: 3194)	SEQ ID NO: 3197)	O ID NO: 3201)	Q ID NO: 3224)	3Q ID NO: 2195)	(ID NO: 3207)	DILPDYDFWNPNEDASSLDT (SEQ ID NO: 3133)
DTTDY (SEQ ID NO: 2203)	SNWGGDAFDI (SEO ID NO: 3202)	LAFDI (SEO ID NO: 3174)	DTTDY (SEO ID NO: 2203)	PAASSRGPKDAFD	LSGDS (SEO ID NO: 3122)	EGVAAGEDY (SEO ID NO: 3123)	FVLDY (SEO ID NO: 2210)	GNGKDV (SEO ID NO: 3135)	EGVAAGEDY (SEO ID NO: 3	DLDFDY (SEO ID NO: 2208)	VNDIVVVDIMDV (SEO ID NO: 3143)	EKRGSRRVFDI (SEO ID NO: 3093)	LSNRNDNLRLDY (SEO ID NO: 3106)	FVLDY (SEQ ID NO: 2210)	TWATNIFDM (SEQ ID NO: 3152)	FDLDY (SEQ ID NO: 3167)	VEWEDIVVGSAFD	GGDMTTVTTDY (SEQ ID NO: 3177)	ADYSNDYYMDV (SEO ID NO: 3166)	EGVAAGEDY (SEQ ID NO: 3123)	GPIYYFDGSAYEG	MNADAFEI (SEO ID NO: 3223)	PAASSRGPKDAFDI (SEQ ID NO: 3129)	DSRPTNRAFHY (SEQ ID NO: 3110)	LHCTGGSCGF (SEQ ID NO: 3186)	VRDDSAGFDY (SEQ ID NO: 3173)	VLVRGQYRGMDL (SEQ ID NO: 3138)	VDYTDYEMGAFD	DRIAAAGGDAFDI (SEQ ID NO: 3194)	DLYKNGYALFDS (SEQ ID NO: 3197	DEYSSLYMDV (SEQ ID NO: 3201	FGAGRLYDDY (SEQ ID NO: 3224)	DNGGGTIGFDY (SEQ ID NO: 2195	DQGIETANDY (SEQ ID NO: 3207)	DILPDYDFWNPNE
99 - 103	99 - 108	99 - 103	99 - 103	99 - 112		•		99 - 104			99 - 109	99 - 109	99 - 110	99 - 103	99 - 107	99 - 103	99-112	99 - 109	99 - 109	99 - 107	99 - 116	98 - 105	99-112	99 - 109	101 - 110	99 - 108	99-110	99-111	99 - 111	99 - 110	99 - 108	801 - 66		99 - 108	99 - 118
99 - 09	20 - 66	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	99 - 09	50 - 66	50 - 66	50 - 66	20 - 66	50 - 66	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	SO - 65	99 - 09	20 - 66	50 - 68	20 - 66	20 - 66	20 - 66	20 - 66	99 - 09	20 - 66	20 - 66	20 - 66	20 - 66	99 - 0ġ
26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35	26 - 35		26 - 35	26 - 35	26 - 35	26-35	26 - 35
1 - 114	1 - 119	1-114	1-114	1 - 123	1-114	1 - 118	1-114	1 - 115	1 - 118	1-115	1 - 120	1 - 120	1 - 121	1 - 114	1 - 118	1 - 114	1 - 123	1 - 120	1 - 120	1-118	1 - 127	1 - 116	1 - 123	1 - 120	1 - 121	1-119	1 - 121	1 - 122	1 - 122	1 - 121	1 - 119	1-119	1 - 120	1-119	1 - 129
219 - 229	221 - 231	220 - 229	219 - 229	228 - 238	220 - 229	223 - 232	220 - 229	219 - 227	222 - 230	221 - 230	222 - 232	222 - 232	223 - 233		220 - 230		227 - 235				229 - 239	218 - 227		•						•		•		1	236 - 245
180 - 186	•	181 - 187	180 - 186	189 - 195	181 - 187	184 - 190	181 - 187	180 - 186	183 - 189	182 - 188	183 - 189	183 - 189	184 - 190	•	•		188 - 194	٠.		189 - 195	961 - 061	179 - 185	186 - 192			187 - 193	187 - 193	190 - 196	961 - 061	188 - 194	187 - 193			•	197 - 203
152 - 164		1	52 - 164	161 - 173	52 - 165	56 - 168	52 - 165	•	57 - 167	53 - 166	57 - 167	1	٠	•	1	,	•	1	7	-173	- 174	- 163	- 170	- 169	- 172	- 171	- 171	- 174	- 174	- 172	- 171	- 171	- 172		- 181
	- 242	- 240	- 240	- 249	- 240	- 243	- 240	- 238	-241	- 241	- 243		- 244	-240	-241	- 240	- 246	- 243		- 249	_	- 238		- 244	- 248	- 248	-247	- 250	- 220	- 246	- 247	- 246		- 246	- 256
2045				****						•			•	•			_	-	_	_	-	_	-		_	_	_ '	- .							•
1081H03	1081H04	1081H06	I081H08	I082A02	· I082A04	I082A08	I082A11	I082B06	I082B09	I082B12	I082C01	1082C05	I082C08	I082D02	I082E05	1082E06	I082E07	I082F11	1082G07	1082G10	1082G11	1082H04	I082H09	I083A06	I083A09	I083A11	1083B03	1083805	1083B06	1083B10	1083C01	1083C02	1083C07	1083C12	1083D04

				•
DEQMVRGVFIANPPIYNYYGMDV (SEQ ID NO: 3154) DADEGLVEAETTNWFDS (SEQ ID NO: 3126) ATKSYDILTRMYYYHMDV (SEQ ID NO: 2748) DRTRMDV (SEQ ID NO: 3182) VGIKAAAVDNFEY (SEQ ID NO: 2197) DEIYNDAFDY (SEQ ID NO: 3105) DGDISDSPINNONYAMDI (SEO ID NO: 3101)		VDYTDYEMGAFDL (SEQ ID NO: 3172) SVAGRGNFDY (SEQ ID NO: 3208) ERGGRDGDYALDF (SEQ ID NO: 3148) EGGGDA YDVAPYYFDY (SEQ ID NO: 2204) DPFDY (SEQ ID NO: 3134) ALLGLPSDFSYYVDY (SEQ ID NO: 3159)	EGEGLICATIVAET TEDT (SEQ ID NO: 3100) TDYGGFDY (SEQ ID NO: 3092) GGVGDSRGYFDP (SEQ ID NO: 3162) DTTDY (SEQ ID NO: 2203) BTTDY (SEQ ID NO: 2203) ESLTGDAFDI (SEQ ID NO: 3116) SPLHFSDAFDI (SEQ ID NO: 3120) DTTDY (SEQ ID NO: 3120) EVGGAFDI (SEQ ID NO: 3157)	DTTDY (SEQ ID NO: 2203) ESLTGDAFDI (SEQ ID NO: 3116)
DFQMVRG DADEGLVI ATKSYDIL DRTRMDV VGIKAAAN DEIYNDAF	RGGTSENY RGGTSENY DYPHNAFI DVRSDRFV STLEVGAT SDDWGAY ERGGRDGI ELVGAPGC	VDYTDYEI SVAGRGNI ERGGRDGI EGGGDAY DPFDY (SE ALLGLPSD	TDYGGFDY TDYGGFDY GGVGDSR DTTDY (SE ESLTGDAF SPLHFSDA DTTDY (SE	DTTDY (SEQ ID SELTGDAFDI (S
99 - 121 99 - 115 102 - 119 99 - 105 99 - 111 99 - 116	99 - 111 99 - 107 99 - 118 99 - 110 99 - 107	99 - 111 101 - 110 99 - 111 99 - 103 99 - 113	99 - 114 99 - 110 99 - 110 99 - 103 99 - 108 99 - 109 99 - 109	99 - 103 99 - 103 99 - 103 99 - 103 99 - 103 99 - 103
50 - 66 50 - 66 52 - 69 50 - 66 50 - 66 50 - 66	50 - 50 - 50 - 50 - 50 - 50 - 50 - 50 -	50 - 68 50 - 68 50 - 68 50 - 68 50 - 68	50-50-50-50-50-50-50-50-50-50-50-50-50-5	50 - 66 50 - 66 50 - 66 50 - 66 50 - 66 50 - 66
26 - 35 26 - 35 26 - 37 26 - 35 26 - 35 26 - 35	50 - 50 - 50 - 50 - 50 - 50 - 50 - 50 -	26,26,26	7 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	26-35 26-35 26-35 26-35 26-35 26-35 26-35
1-132 1-126 1-130 1-116 1-122 1-119		1 1 1 1 1 1	1 - 120 1 - 114 1 - 114 1 - 114 1 - 116 1 - 120 1 - 114	1 - 114 1 - 114 1 - 114 1 - 114 1 - 114 1 - 114
243 - 251 233 - 243 236 - 247 221 - 231 228 - 238 226 - 237			224 - 242 222 - 232 226 - 236 218 - 226 219 - 229 221 - 231 222 - 232 219 - 229	218 - 226 218 - 226 216 - 226 219 - 229 218 - 226 218 - 226 225 - 234
204 - 210 194 - 200 197 - 203 182 - 188 189 - 195 187 - 193	188 - 194 186 - 192 197 - 203 188 - 194 186 - 192 190 - 196	190 - 196 189 - 195 190 - 196 193 - 199 182 - 188 192 - 198	193 - 199 183 - 189 187 - 193 179 - 185 180 - 186 182 - 188 183 - 189 180 - 186	179 - 185 177 - 183 177 - 183 180 - 186 179 - 185 179 - 185
173 - 188 165 - 178 169 - 181 156 - 166 161 - 173 158 - 171			164 - 177 157 - 167 161 - 171 153 - 163 152 - 164 157 - 167 152 - 164 155 - 168	153 - 163 153 - 163 151 - 161 152 - 164 153 - 163 153 - 163
148 - 262 142 - 254 146 - 258 132 - 242 138 - 249 135 - 248 143 - 255	248 248 248 247 247 250	-250 -249 -253 -252	141 - 253 133 - 243 137 - 247 130 - 237 135 - 242 136 - 243 130 - 240 133 - 243	130 - 237 130 - 237 130 - 237 130 - 240 130 - 237 130 - 237
2081 2083 2083 2084 2085 2085	2088 2089 2090 2091 2093 2093	2095 2096 2097 2098 2100	2102 2103 2104 2105 2107 2108 2108	2110 2111 2112 2113 2114 2115 2115
1083D07 1083D10 1083D10 1083D12 1083E02 1083E03	1083E08 1083E12 1083F02 1083F04 1083F06 1083F08 1083F11	1083G04 1083G05 1083G06 1083G08 1083G11	1083H04 1083H05 1083H07 1084A03 1084A08 1084B08 1084C02 1084D03	1084E01 1084E06 1084E10 1084E12 1084F04 1084F07

•		•									
DTTDY (SEQ ID NO: 2203)	-	O	VGIKAAAVDNFEY (SEO ID NO: 2197)	LGRNYTSSWSLDY (SEO ID NO: 3181)	VGIKAAAVDNFEY (SEO ID NO: 2197)	GGRYGYYYDGTGYYDAFDI (SEO ID NO: 3226)			DNGGGTIGFDY (SEQ ID NO: 2195)	DNGGGTIGFDY (SEO ID NO: 2195)	VRQQIADPPRSFFDP (SEQ ID NO: 3144)
99 - 103	99 - 103	99 - 118	99 - 111		99 - 111	99 - 117		99-113		99 - 109	99 - 113
50 - 66	50 - 66	99 - 09	50 - 66	99 - 09	50 - 66	50 - 66	99-09	99 - 09	50 - 66	50 - 66	26-35 50-66
26 - 35	1-114 26-35 50-66	26 - 35	26-35	26 - 35	26 - 35	26 - 35	26-35	26 - 35	26 - 35	26 - 35	
1-114	1-114	1 - 129	1 - 122	1 - 122	1 - 122	1 - 128	1 - 120	1 - 124	1 - 120	1-120	1 - 124
219 - 229	218 - 226	235 - 245	228 - 238	227 - 237	228 - 238	234 - 244	227 - 236	230 - 240	227 - 236	227 - 236	230 - 240
80 - 186	79-185	96 - 202	89 - 195	88 - 194	89 - 195	95 - 201	88 - 194	91 - 197	88 - 194	88 - 194	191 - 197
152 - 164	153 - 163	168 - 180	161 - 173	162 - 172	161 - 173	167 - 179	159 - 172	163 - 175	159 - 172	159 - 172	163 - 175
130 - 240	130 - 237	145 - 256	138 - 249 161 - 173 1	138 - 248	138 - 249	144 - 255	136 - 247	140 - 251	136 - 247	136 - 247	140 - 251
			2120								
1084G12	1084H02	1099B05	1099G09	1099H01	90H660I	1099H08	1100A01	1100A10	1100B03	1100B04	1100C03

	ING TO A DEPOSITED MICROORGANISM ER BIOLOGICAL MATERIAL
	(PCT Rule 13bis)
A. The indications made below relate to the deposite description on page 20, paragraph 58.	ed microorganism or other biological material referred to in the
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution: American T	ype Culture Collection
Address of depositary institution (including 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America	postal code and country)
Date of deposit 22 October 1996	Accession Number 97768
C. ADDITIONAL INDICATIONS (leave blank if no	ot applicable) This information is continued on an additional sheet
D. DESIGNATED STATES FOR WHICH INDIC	CATIONS ARE MADE (if the indications are not for all designated States)
until the publication of the mention of the grant of the E	atent is sought a sample of the deposited microorganism will be made available European patent or until the date on which the application has been refused or sue of such a sample to an expert nominated by the person requesting the Continued on additional sheets
E. SEPARATE FURNISHING OF INDICATION	NS (leave blank if not applicable)
The indications listed below will be submitted to the intern Number of Deposit")	national Bureau later (specify the general nature of the indications e.g., "Accession
For receiving Office use only	For International Bureau use only
☐ This sheet was received with the international application	on
Authorized officer	Authorized officer
evised Form PCT/PO/134 (Ignuary 2001)	

ATCC Deposit No.: 97768

CANADA

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: 97768

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

PCT/US01/19110 WO 02/02641

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

OR OTHER BIOLOGICAL MATERIAL			
(I	PCT Rule 13bis)		
A. The indications made below relate to the deposited mi description on page 20, paragraph 59.	croorganism or other biological material referred to in the		
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet		
Name of depositary institution: American Type	Culture Collection		
Address of depositary institution (including posi 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America	tal code and country)		
Date of deposit 10 December 1998	Accession Number 203518		
C. ADDITIONAL INDICATIONS (leave blank if not app	plicable) This information is continued on an additional sheet		
•			
D DESIGNATED STATES FOR WHICH INDICAT	IONS ARE MADE (if the indications are not for all designated States)		
until the publication of the mention of the grant of the Europ	is sought a sample of the deposited microorganism will be made available bean patent or until the date on which the application has been refused or of such a sample to an expert nominated by the person requesting the Continued on additional sheets		
E. SEPARATE FURNISHING OF INDICATIONS (let	ave blank if not applicable)		
The indications listed below will be submitted to the internation Number of Deposit")	nal Bureau later (specify the general nature of the indications e.g., "Accession		
For receiving Office use only	For International Bureau use only		
☐ This sheet was received with the international application	☐ This sheet was received by the International Bureau on:		
Authorized officer	Authorized officer		
Revised Form PCT/RO/134 (January 2001)	Pctro134ep.solli		

ATCC Deposit No.: 203518

CANADA

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: 203518

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

PCT/US01/19110 WO 02/02641

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

OR OTHER BIOLOGICAL MATERIAL					
(PCT Rule 13bis)					
A. The indications made below relate to the deposited microorganism or other biological material referred to in the description on page 145, Table 2.					
B. IDENTIFICATION OF DEPOSIT	3. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet				
Name of depositary institution: American Type Culture Collection					
Address of depositary institution (including postal code and country) 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America					
Date of deposit 27 March 2001		Accession Number PTA-3238			
C. ADDITIONAL INDICATIONS (leave blank ij	f not applic	able)	This information is continued on an addition	nal sheet	
	 ;				
D. DESIGNATED STATES FOR WHICH IND	ICATIO	NS ARE MAI	DE (if the indications are not for all designated St	ates)	
Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused or withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). Continued on additional sheets					
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)					
The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Accession Number of Deposit")					
For receiving Office use only			For International Bureau use only		
☐ This sheet was received with the international applic	cation	☐ This sheet was received by the International Bureau on:			
Authorized officer	Authorized officer				

ATCC Deposit No.: PTA-3238

CANADA

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: PTA-3238

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM OR OTHER BIOLOGICAL MATERIAL

OR OTHE	R BIOLOGICAL MATERIAL
	(PCT Rule 13bis)
A. The indications made below relate to the deposited description on page 145, Table 2.	microorganism or other biological material referred to in the
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution: American Ty	pe Culture Collection
Address of depositary institution (including p 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America	ostal code and country)
Date of deposit 27 March 2001	Accession Number PTA-3239
C. ADDITIONAL INDICATIONS (leave blank if not	applicable) This information is continued on an additional sheet
D. DESIGNATED STATES FOR WHICH INDICA	ATIONS ARE MADE (if the indications are not for all designated States)
until the publication of the mention of the grant of the Eur	ant is sought a sample of the deposited microorganism will be made available ropean patent or until the date on which the application has been refused or such a sample to an expert nominated by the person requesting the Continued on additional sheets
E. SEPARATE FURNISHING OF INDICATIONS	(leave blank if not applicable)
The indications listed below will be submitted to the internate Number of Deposit")	ional Bureau later (specify the general nature of the indications e.g., "Accession
For receiving Office use only	For International Bureau use only
☐ This sheet was received with the international application	☐ This sheet was received by the International Bureau on:
Authorized officer	Authorized officer
evised Form PCT/RO/134 (January 2001)	Petrol 34ep.sollis

359

ATCC Deposit No.: PTA-3239

CANADA

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: PTA-3239

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM OR OTHER BIOLOGICAL MATERIAL

OR OTHER B	IOLOGICAL MATERIAL				
(P)	CT Rule 13bis)				
A. The indications made below relate to the deposited mid description on page 145, Table 2.	croorganism or other biological material referred to in the				
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet					
Name of depositary institution: American Type	Culture Collection				
Address of depositary institution (including posta 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America	al code and country)				
Date of deposit	Accession Number				
27 March 2001	PTA-3240				
C. ADDITIONAL INDICATIONS (leave blank if not app	olicable) This information is continued on an additional sheet				
D DESIGNATED STATES FOR WHICH INDICATE	ONS ARE MADE (if the indications are not for all designated States)				
until the publication of the mention of the grant of the Europe	s sought a sample of the deposited microorganism will be made available ean patent or until the date on which the application has been refused or f such a sample to an expert nominated by the person requesting the Continued on additional sheets				
E. SEPARATE FURNISHING OF INDICATIONS (lea	we blank if not applicable)				
	al Bureau later (specify the general nature of the indications e.g., "Accession				
For receiving Office use only	For International Bureau use only				
☐ This sheet was received with the international application	☐ This sheet was received by the International Bureau on:				
Authorized officer	Authorized officer				
Perisad Form PCT/PO/124 (Innuary 2001)	Potmi 3den soll				

ATCC Deposit No.: PTA-3240

CANADA

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: PTA-3240

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

OR OTHER BIOLOGICAL MATERIAL					
(PCT Rule 13bis)					
A. The indications made below relate to the deposited microorganism or other biological material referred to in the description on page 145, Table 2.					
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet				
Name of depositary institution: American Type Culture Collection					
Address of depositary institution (including postal 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America	l code and country)				
Date of deposit	Accession Number				
27 March 2001	PTA-3241				
C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet					
	•				
D. DESIGNATED STATES FOR WHICH INDICATION	ONS ARE MADE (if the indications are not for all designated States)				
until the publication of the mention of the grant of the Europe	sought a sample of the deposited microorganism will be made available an patent or until the date on which the application has been refused or such a sample to an expert nominated by the person requesting the Continued on additional sheets				
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)					
The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Accession Number of Deposit")					
For receiving Office use only	For International Bureau use only				
☐ This sheet was received with the international application	☐ This sheet was received by the International Bureau on:				
Authorized officer	Authorized officer				

365

ATCC Deposit No.: PTA-3241

CANADA ·

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: PTA-3241

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM OR OTHER BIOLOGICAL MATERIAL

Address of depositary institution (including postal code and country) 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America Date of deposit 27 March 2001 PTA-3242 C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet
Name of depositary institution: American Type Culture Collection Address of depositary institution (including postal code and country) 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America Date of deposit 27 March 2001 PTA-3242 C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States) Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made availantil the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). Continued on additional sheets E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access")
Manassas, Virginia 20110-2209 United States of America Date of deposit 27 March 2001 Accession Number PTA-3242 C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States) Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made availuntil the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access
Manassas, Virginia 20110-2209 United States of America Date of deposit 27 March 2001 PTA-3242 C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States) Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made availuntil the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access")
C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States) Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made availuntil the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). Continued on additional sheets E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States) Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). Continued on additional sheets E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access
Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). Continued on additional sheets E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access")
Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). Continued on additional sheets E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access")
Europe In respect of those designations in which a European Patent is sought a sample of the deposited microorganism will be made available until the publication of the mention of the grant of the European patent or until the date on which the application has been refused withdrawn or is deemed to be withdrawn, only by the issue of such a sample to an expert nominated by the person requesting the sample (Rule 28(4) EPC). Continued on additional sheets E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access")
The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Access
For receiving Office use only For International Bureau use only
☐ This sheet was received with the international application ☐ This sheet was received by the International Bureau on:
Authorized officer Authorized officer

ATCC Deposit No.: PTA-3242

CANADA

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: PTA-3242

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

OR (OTHER BI	OLOGICAL 1	MATERIAL			
	(PC	T Rule 13bis))			
A. The indications made below relate to the dep description on page 145, Table 2.	posited mici	roorganism or o	other biological material referred to in the			
B. IDENTIFICATION OF DEPOSIT	T Further deposits are identified on an additional sheet					
Name of depositary institution: Americ	an Type (Culture Colle	ection			
Address of depositary institution <i>(includ</i> 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America	ling posta	l code and c	country)			
Date of deposit 27 March 2001		Accession Nu	umber PTA-3243			
C. ADDITIONAL INDICATIONS (leave blan	k if not appli	icable)	This information is continued on an additional sheet			
		*	*			
D. DESIGNATED STATES FOR WHICH I	NDICATIO	NS ARE MA	ADE (if the indications are not for all designated States)			
until the publication of the mention of the grant of	the Europea	en patent or unti	le of the deposited microorganism will be made available til the date on which the application has been refused or to an expert nominated by the person requesting the Continued on additional sheets			
E. SEPARATE FURNISHING OF INDICAT	TONS (leave	e blank if not applical	able)			
The indications listed below will be submitted to the indications listed below will be submitted to the indications of Deposit")	international	Bureau later (sp	pecify the general nature of the indications e.g., "Accession			
For receiving Office use only			For International Bureau use only			
☐ This sheet was received with the international application		☐ This sheet was received by the International Bureau on:				
thorized officer		Authorized officer				
Levised Form PCT/RO/134 (January 2001)			Poten 124 on an			

ATCC Deposit No.: PTA-3243

CANADA

The applicant requests that, until either a Canadian patent has been issued on the basis of an application or the application has been refused, or is abandoned and no longer subject to reinstatement, or is withdrawn, the Commissioner of Patents only authorizes the furnishing of a sample of the deposited biological material referred to in the application to an independent expert nominated by the Commissioner, the applicant must, by a written statement, inform the International Bureau accordingly before completion of technical preparations for publication of the international application.

NORWAY

The applicant hereby requests that the application has been laid open to public inspection (by the Norwegian Patent Office), or has been finally decided upon by the Norwegian Patent Office without having been laid open inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Norwegian Patent Office not later than at the time when the application is made available to the public under Sections 22 and 33(3) of the Norwegian Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on the list of recognized experts drawn up by the Norwegian Patent Office or any person approved by the applicant in the individual case.

AUSTRALIA

The applicant hereby gives notice that the furnishing of a sample of a microorganism shall only be effected prior to the grant of a patent, or prior to the lapsing, refusal or withdrawal of the application, to a person who is a skilled addressee without an interest in the invention (Regulation 3.25(3) of the Australian Patents Regulations).

FINLAND

The applicant hereby requests that, until the application has been laid open to public inspection (by the National Board of Patents and Regulations), or has been finally decided upon by the National Board of Patents and Registration without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art.

UNITED KINGDOM

The applicant hereby requests that the furnishing of a sample of a microorganism shall only be made available to an expert. The request to this effect must be filed by the applicant with the International Bureau before the completion of the technical preparations for the international publication of the application.

ATCC Deposit No.: PTA-3243

DENMARK

The applicant hereby requests that, until the application has been laid open to public inspection (by the Danish Patent Office), or has been finally decided upon by the Danish Patent office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the Danish Patent Office not later that at the time when the application is made available to the public under Sections 22 and 33(3) of the Danish Patents Act. If such a request has been filed by the applicant, any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Danish Patent Office or any person by the applicant in the individual case.

SWEDEN

The applicant hereby requests that, until the application has been laid open to public inspection (by the Swedish Patent Office), or has been finally decided upon by the Swedish Patent Office without having been laid open to public inspection, the furnishing of a sample shall only be effected to an expert in the art. The request to this effect shall be filed by the applicant with the International Bureau before the expiration of 16 months from the priority date (preferably on the Form PCT/RO/134 reproduced in annex Z of Volume I of the PCT Applicant's Guide). If such a request has been filed by the applicant any request made by a third party for the furnishing of a sample shall indicate the expert to be used. That expert may be any person entered on a list of recognized experts drawn up by the Swedish Patent Office or any person approved by a applicant in the individual case.

NETHERLANDS

The applicant hereby requests that until the date of a grant of a Netherlands patent or until the date on which the application is refused or withdrawn or lapsed, the microorganism shall be made available as provided in the 31F(1) of the Patent Rules only by the issue of a sample to an expert. The request to this effect must be furnished by the applicant with the Netherlands Industrial Property Office before the date on which the application is made available to the public under Section 22C or Section 25 of the Patents Act of the Kingdom of the Netherlands, whichever of the two dates occurs earlier.

WHAT IS CLAIMED IS:

1. An antibody that immunospecifically binds to BLyS comprising a first amino acid sequence at least 95% identical to an second amino acid sequence selected from the group consisting of:

- (a) an amino acid sequence comprising the amino acid sequence of a VHCDR of any one of the scFvs of SEQ ID NOS:1 through 2128; and
- (b) an amino acid sequence comprising the amino acid sequence of a VLCDR of any one of the scFvs of SEQ ID NOS:1 through 2128.
- The antibody of claim 1, wherein the second amino acid sequence consists of the amino acid sequence of a VHCDR3 of any one of the scFvs of SEQ ID NOS:2129 through 3227.
- 3. The antibody of claim 1, wherein the second amino acid sequence consists of the amino acid sequence of a VH domain of any one of the scFvs of SEQ ID NOS:1 through 2128.
- 4. The antibody of claim 3 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1 through 1562.
- 5. The antibody of claim 4 in which said antibody immunospecifically binds to both the soluble form and membrane-bound form of BLyS.
- 6. The antibody of claim 4 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1 through 46 and 321 through 329.

7. The antibody of claim 6 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 2, 9, and 327.

- 8. The antibody of claim 4 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 834 through 872.
- 9. The antibody of claim 3 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1563 through 1880.
- 10. The antibody of claim 9 in which, and in which said antibody immunospecifically binds to the soluble form of BLyS.
- 11. The antibody of claim 9 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1563 through 1569.
- 12. The antibody of claim 9 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1570 through 1595.
- 13. The antibody of claim 3 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1881 through 2128.
- 14. The antibody of claim 13 in which said antibody immunospecifically binds to the membrane-bound form of BLyS.
- 15. The antibody of claim 13 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1881 through 1885.
- 16. The antibody of claim 13 in which said VH domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 1886 through 1908.

17. The antibody of claim 1, wherein the second amino acid sequence consists of the amino acid sequence of a VL domain of any one of the scFvs of SEQ ID NOS:1 through 2128.

- 18. The antibody of claim 17 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1 through 1562.
- 19. The antibody of claim 18 in which said antibody immunospecifically binds to both the soluble form and membrane-bound form of BLyS.
- 20. The antibody of claim 18 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1 through 46 and 321 through 329.
- 21. The antibody of claim 20 in which said VL domain consists of the amino acid sequence of the VH domain of any one of the scFvs of SEQ ID NOS: 2, 9, and 327.
- 22. The antibody of claim 18 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 834 through 872.
- 23. The antibody of claim 17 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1563 through 1880.
- 24. The antibody of claim 23 said antibody immunospecifically binds to the soluble form of BLyS.
- 25. The antibody of claim 23 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1563 through 1569.

26. The antibody of claim 23 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1570 through 1595.

- 27. The antibody of claim 17 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1881 through 2128.
- 28. The antibody of claim 27 in which said antibody immunospecifically binds to the membrane-bound form of BLyS.
- 29. The antibody of claim 27 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1881 through 1885.
- 30. The antibody of claim 27 in which said VL domain consists of the amino acid sequence of the VL domain of any one of the scFvs of SEQ ID NOS: 1886 through 1908.
- 31. The antibody of claim 3, which also comprises an amino acid sequence at least 95% identical to the amino acid sequence of a VL domain of any one of the scFvs of SEQ ID NOS:1 through 2128.
- 32. The antibody of claim 31, wherein the VH and VL domains are from the same scFv.
 - 33. The antibody of claim 32, wherein the scFv is the scFv of SEQ ID NO:2.
 - 34. The antibody of claim 32, wherein the scFv is the scFv of SEQ ID NO:9.
 - 35. The antibody of claim 32, wherein the scFv is the scFv of SEQ ID NO:327.
- 36. The antibody of claim 1 wherein the first amino acid sequence is identical to the second amino acid sequence.

37. The antibody of claim 36 wherein the second amino acid sequence consists of the amino acid sequence of a VH domain of any one of the scFvs of SEQ ID NOS:1 through 2128.

- 38. The antibody of claim 36 wherein the second amino acid sequence consists of the amino acid sequence of a VL domain of any one of the scFvs of SEQ ID NOS:1 through 2128.
- 39. The antibody of claim 37 which also comprises an amino acid sequence 100% identical to the amino acid sequence of a VL domain of any one of the scFvs of SEQ ID NOS:1 through 2128.
 - 40. The antibody of claim 39, wherein the scFv is the scFv of SEQ ID NO:2.
 - 41. The antibody of claim 39, wherein the scFv is the scFv of SEQ ID NO:9.
 - 42. The antibody of claim 39, wherein the scFv is the scFv of SEQ ID NO:327.
 - 43. The antibody of claim 1 through wherein the BLyS is a BLyS homotrimer.
- 44. The antibody of claim 43, wherein the individual protein components of the BLyS homotrimer consist of the mature form of BLyS.
- 45. The antibody of any one of claims 1 through 44, wherein the BLyS is a BLyS heterotrimer.
- 46. The antibody of claim 45, wherein the BLyS heterotrimer comprises at least one BLyS polypeptide and at least one APRIL polypeptide.

- 47. The antibody of claim 46, wherein the BLyS polypeptide consists of the mature form of BLyS and the APRIL polypeptide consists of the mature form of APRIL.
- 48. The antibody of any one of claims 1 through 47, wherein the antibody is selected from the group consisting of:
 - (a) a whole immunoglobulin molecule;
 - (b) an scFv;
 - (c) a monoclonal antibody;
 - (d) a human antibody;
 - (e) a chimeric antibody;
 - (f) a humanized antibody;
 - (g) a Fab fragment;
 - (h) an Fab' fragment;
 - (i) an F(ab')2;
 - (j) an Fv; and
 - (k) a disulfide linked Fv.
- 49. The antibody of claim 3 or 37, which also comprises a heavy chain immunoglobulin constant domain selected from the group consisting of:
 - (a) a human IgM constant domain;
 - (b) a human IgG1 constant domain;
 - (c) a human IgG2 constant domain;
 - (d) a human IgG3 constant domain;
 - (e) a human IgG4 constant domain; and
 - (f) a human IgA constant domain.

50. The antibody of claim 17 or 38, which also comprises a light chain immunoglobulin constant domain sleeted from the group consisting of:

- (a) a human Ig kappa constant domain;
- (b) a human Ig lambda constant domain.
- 51. The antibody of any one of claims 1 through 50, wherein the antibody has a dissociation constant (K_D) selected from the group consisting of:
 - (a) a dissociation constant (K_D) between 10⁻⁷ M and 10⁻⁸ M;
 - (b) a dissociation constant (K_D) between 10⁻⁸ M and 10⁻⁹ M;
 - (c) a dissociation constant (K_D) between 10⁻⁹ M and 10⁻¹⁰ M;
 - (d) a dissociation constant (K_D) between 10^{-10} M and 10^{-11} M;
 - (e) a dissociation constant (K_D) between 10⁻¹¹ M and 10⁻¹² M; and
 - (f) a dissociation constant (K_D) between 10^{-12} M and 10^{-13} M.
- 52. The antibody of any one of claims 1 through 51, wherein the antibody is conjugated to a detectable label.
 - 53. The antibody of claim 52, wherein the detectable label is a radiolabel.
- 54. The antibody of claim 53, wherein the radiolabel is ¹²⁵I, ¹³¹I, ¹¹¹In, ⁹⁰Y, ⁹⁹Tc, ¹⁷⁷Lu, ¹⁶⁶Ho, or ¹⁵³Sm.
- 55. The antibody of claim 52, wherein the detectable label is an enzyme, a fluorescent label, a luminescent label, or a bioluminescent label.
- 56. The antibody of any one of claims 1 through 51, wherein the antibody is biotinylated.

57. The antibody of any one of claims 1 through 51, wherein the antibody is conjugated to a therapeutic or cytotoxic agent.

- 58. The antibody of claim 57, wherein the therapeutic or cytotoxic agent is selected from the group consisting of:
 - (a) an anti-metabolite,
 - (b) an alkylatingagent;
 - (c) an antibiotic;
 - (d) a growth factor;
 - (e) a cytokine;
 - (f) an anti-angiogenic agent;
 - (g) an anti-mitotic agent;
 - (h) an anthracycline;
 - (i) toxin; and
 - (j) an apoptotic agent.
- 59. An antibody of any one of claims 1 through 58, that neutralizes BLyS or a fragment thereof.
- 60. The antibody of claim 59, that diminishes or abolishes the ability of BLyS or a fragment thereof to bind to its receptor.
 - 61. The antibody of claim 60, wherein the receptor is TACI.
 - 62. The antibody of claim 60, wherein the receptor is BCMA.
- 63. The antibody of claim 59, that diminishes or abolishes the ability of BLyS or a fragment thereof to stimulate B cell proliferation.

64. The antibody of claim 59, that diminishes or abolishes the ability of BLyS or a fragment thereof to stimulate immunoglobulin secretion by B cells.

- 65. An antibody of any one of claims 1 through 58, that enhances the activity of BLyS or a fragment thereof.
- 66. The antibody of claim 65, that increases the ability of BLyS or a fragment thereof to bind to its receptor.
 - 67. The antibody of claim 66, wherein the receptor is TACI.
 - 68. The antibody of claim 66, wherein the receptor is BCMA.
- 69. The antibody of claim 65, that increases the ability of BLyS or a fragment thereof to stimulate B cell proliferation.
- 70. The antibody of claim 65, that increases the ability of BLyS or a fragment thereof to stimulate immunoglobulin secretion by B cells.
- 71. The antibody of any one of claims 1 through 70 covalently linked to a heterologous polypeptide.
- 72. The antibody of claim 71, wherein the heterologous polypeptide is human serum albumin.
- 73. The antibody of any one of claims 1 through 72 in a pharmaceutically acceptable carrier.
 - 74. A kit comprising the antibody of any one of claims 1 through 73.

75. An isolated nucleic acid molecule encoding the antibody of any one of claims 1 through 74.

- 76. A vector comprising the isolated nucleic acid molecule of claim 75.
- 77. The vector of claim 76 which also comprises a nucleotide sequence which regulates the expression of the antibody encoded by the nucleic acid molecule.
 - 78. A host cell comprising the nucleic acid molecule of claim 77.
- 79. A cell line engineered to express the antibody of any one of claims 1 through 78.
- 80. An antibody that binds the same epitope as the antibody of any one of claims 1 through 79.
- 81. An antibody that competitively inhibits the binding of the antibody produced by the cell line having ATCCDeposit Number PTA-3239.
- 82. An antibody that competitively inhibits the binding of the antibody produced by the cell line having ATCCDeposit Number PTA-3240
- 83. An antibody that competitively inhibits the binding of the antibody produced by the cell line having ATCCDeposit Number PTA-3243.
- 84. An second antibody that reduces the binding of the antibody of any one of claims 1 through 83 by a an increment within a percentage range selected from the group consisting of:
 - (a) from 50% up to 60%;

- (b) from 60% up to 70%;
- (c) from 70% up to 80%;
- (d) from 80% up to 90%; and
- (e) from 90% up to 100%.
- 85. An antibody that immunospecifically binds to BLyS, said antibody comprising an amino acid sequence of a VH domain encoded by a nucleotide sequence that hybridizes under stringent conditions to a nucleotide sequence encoding a VH domain of an scFv comprising an amino acid sequence of any one of SEQ ID NOS: 1 to 2128.
- 86. An antibody that immunospecifically binds to BLyS, said antibody comprising an amino acid sequence of a VL domain encoded by a nucleotide sequence that hybridizes under stringent conditions to a nucleotide sequence encoding a VL domain from an scFv comprising an amino acid sequence of any one of SEQ ID NOS: 1 to 2128.
 - 87. A method for detecting aberrant expression of BLyS protein, comprising:
- (a) assaying the level of BLyS expression in a first biological sample of an individual using one or more antibodies of any one of claims 1 through 86; and
- (b) comparing the level of BLyS assayed in biological sample with a standard level of BLyS expression or level of BLyS in a second, normal biological sample;
- (c) wherein an increase or decrease in the assayed level of BLyS in the first biological sample compared to the standard level of BLyS expression or level of BLyS in a second, normal biological sample, is indicative of aberrant expression.
- 88. A method for diagnosing a disease or disorder associated with aberrant BLyS expression or activity, comprising:
- (a) administering to a subject an effective amount of a labeled antibody of any one of claims 52 through 58 that immunospecifically binds to BLyS;

(b) waiting for a time interval following the administering for permitting the labeled antibody of any one of claims 52 through 58 to preferentially concentrate at sites in the subject where BLyS is expressed;

- (c) determining background level; and
- (d) detecting the labeled antibody of any one of claims 52 through 58 in the subject, such that detection of labeled antibody above the background level indicates that the subject has a particular disease or disorder associated with aberrant expression of BLyS.
- 89.A method of treating, preventing or ameliorating a disease or disorder associated with aberrant BLyS expression or activity, comprising administering to an animal in need thereof, the pharmaceutical composition of claim 73 in an amount effective to treat, prevent or ameliorate the disease or disorder.
 - 90. The method of claim 89, wherein the disease or disorder is cancer.
- 91. The method of claim 89, wherein the disease or disorder of the immune system.
- 92. The method of claim 91, wherein the disease or disorder of the immune system is an autoimmune disease or disorder.
- 93. The method of claim 92, wherein the disease or disorder of the immune system is an autoimmune disease or disorder selected from the group consisting of:
 - (a) Systemic Lupus Erythematosus; and
 - (b) Rheumatoid Arthritis.

94. The method of claim 91, wherein the disease or disorder of the immune system is an immunodeficiency.

- 95. The method of claim 92, wherein the disease or disorder of the immune system is an immunodeficiency selected from the group consisting of:
 - (a) Common Variable Immunodeficiency (CVID); and
 - (b) AIDS.
- 96. The method of claim 91, wherein the disease or disorder of the immune system is cancer.

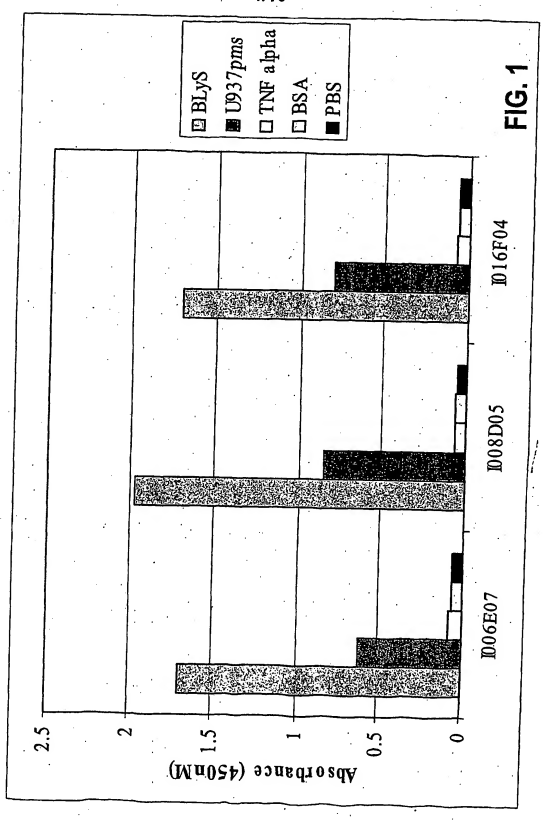
94. The method of claim 91, wherein the disease or disorder of the immune system is an immunodeficiency.

95. The method of claim 92, wherein the disease or disorder of the immune system is an immunodeficiency selected from the group consisting of:

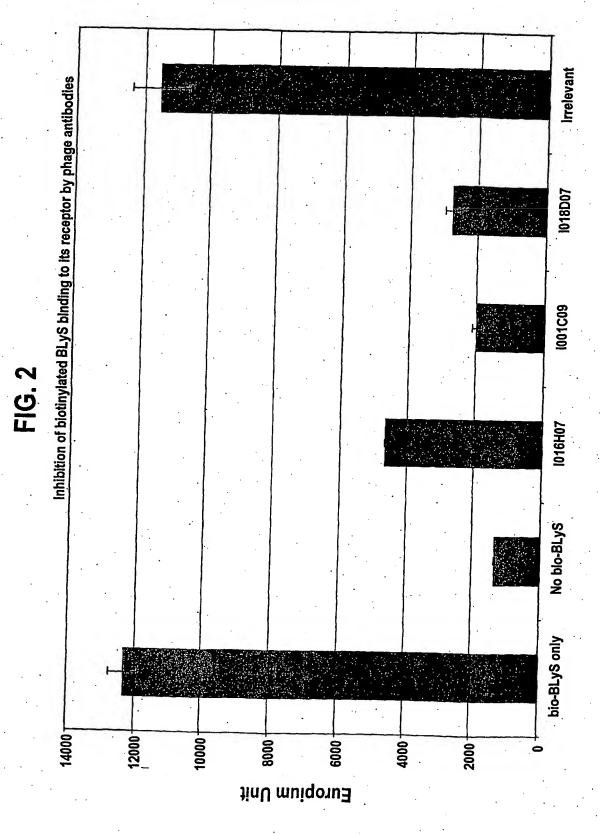
- (a) Common Variable Immunodeficiency (CVID); and
- (b) AIDS.

96. The method of claim 91, wherein the disease or disorder of the immune system is cancer.

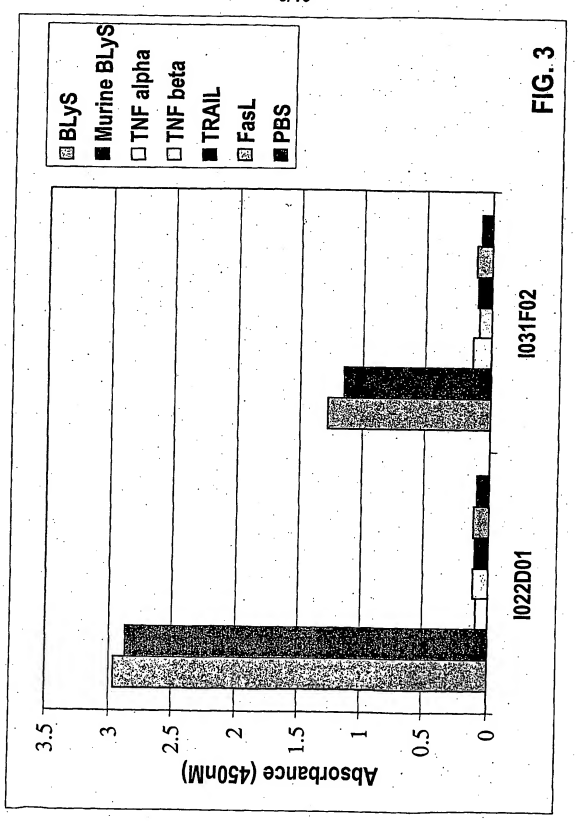
1/16



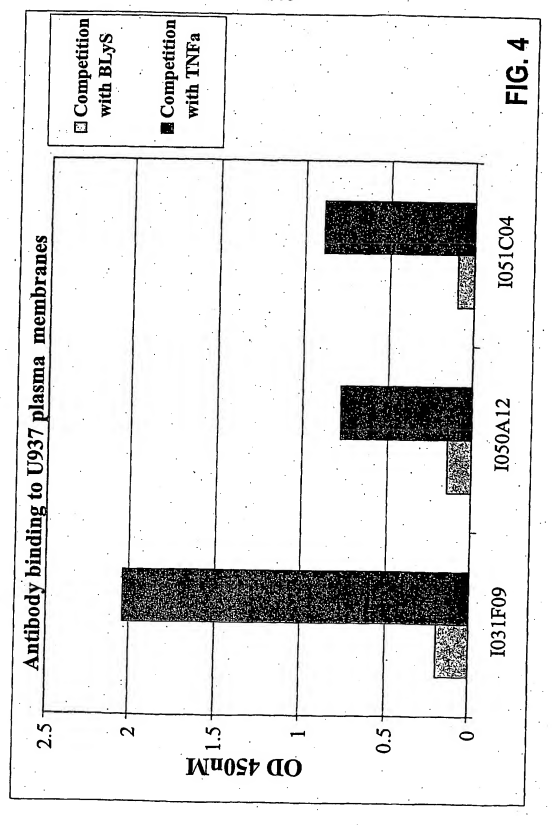


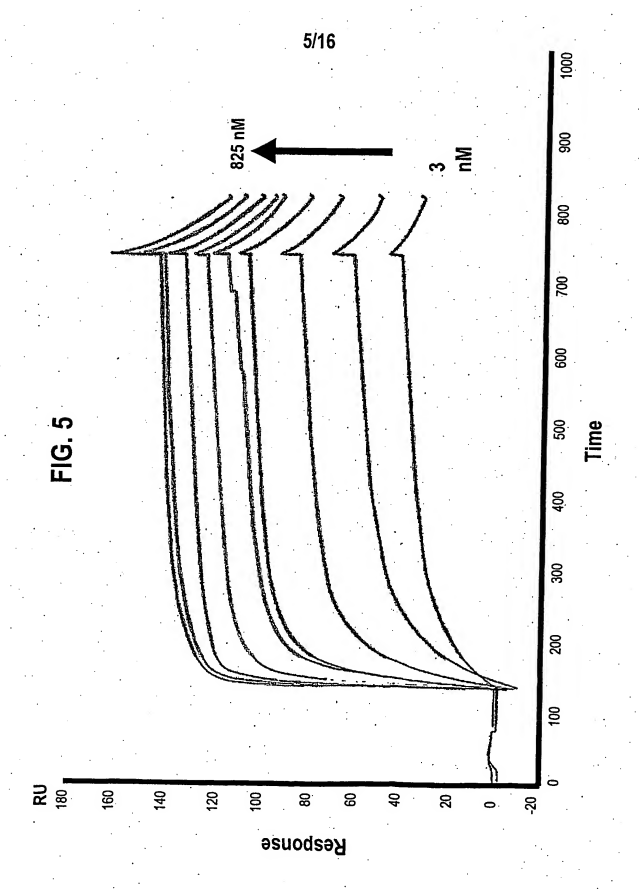




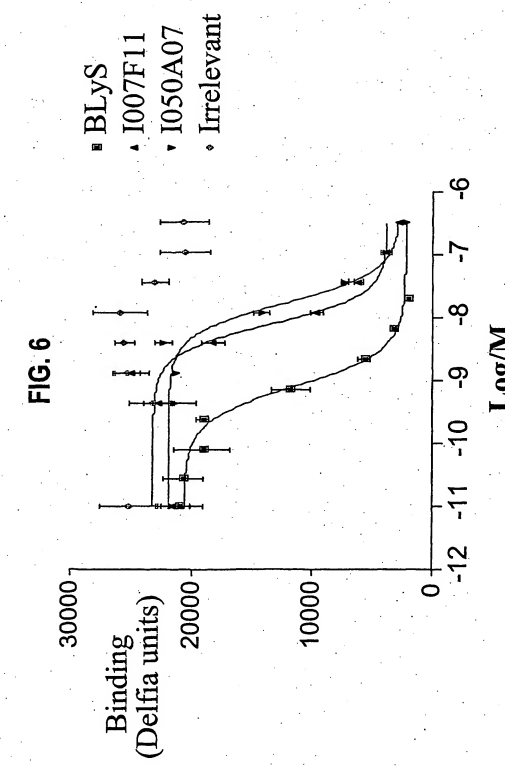




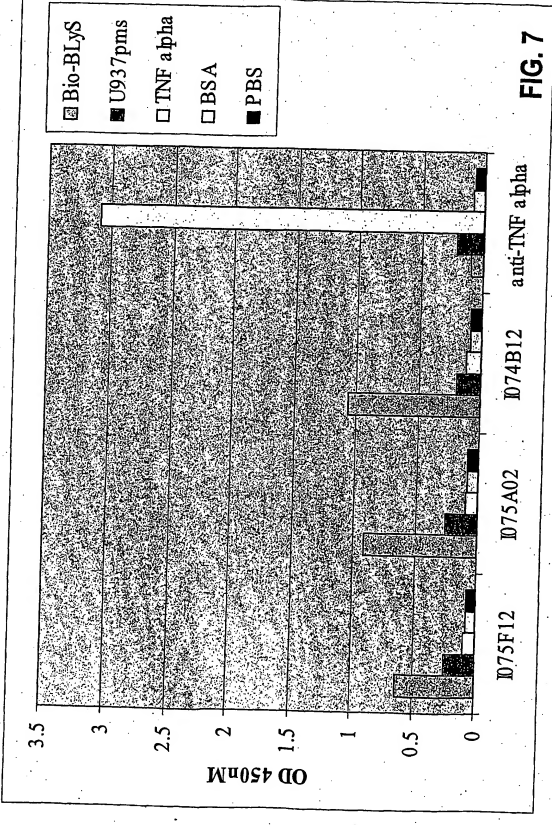


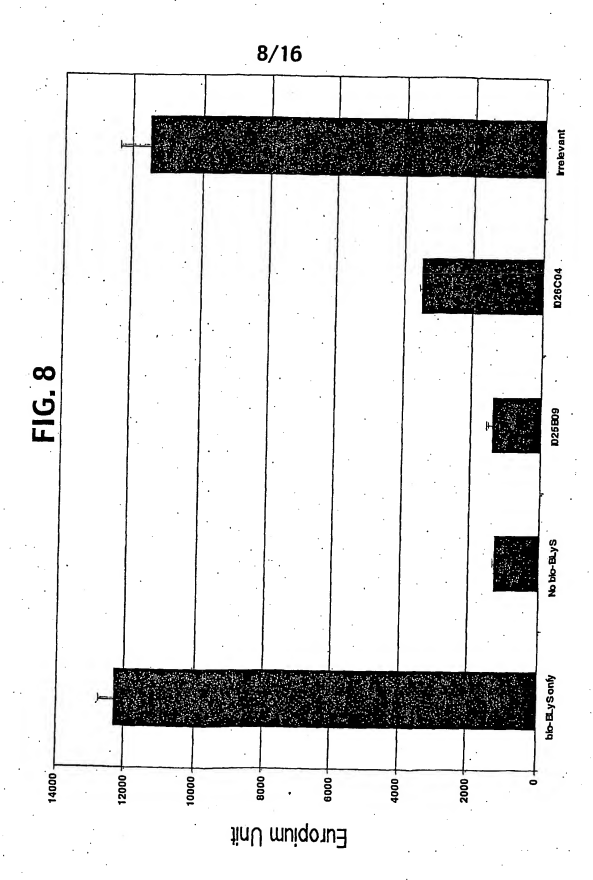


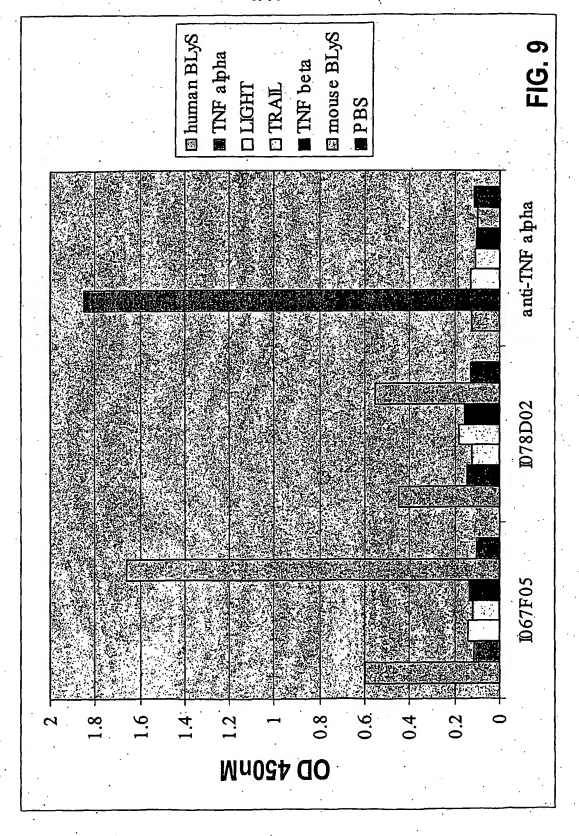


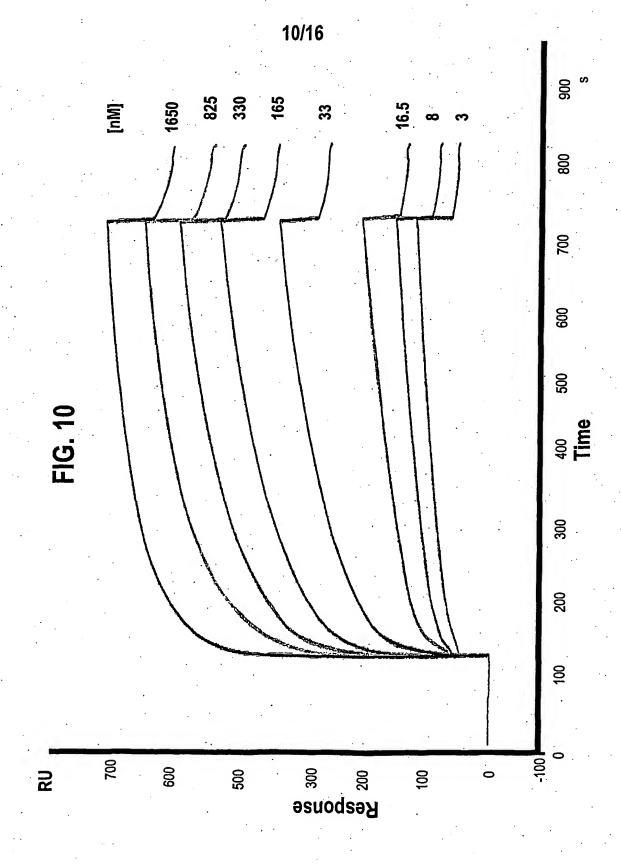


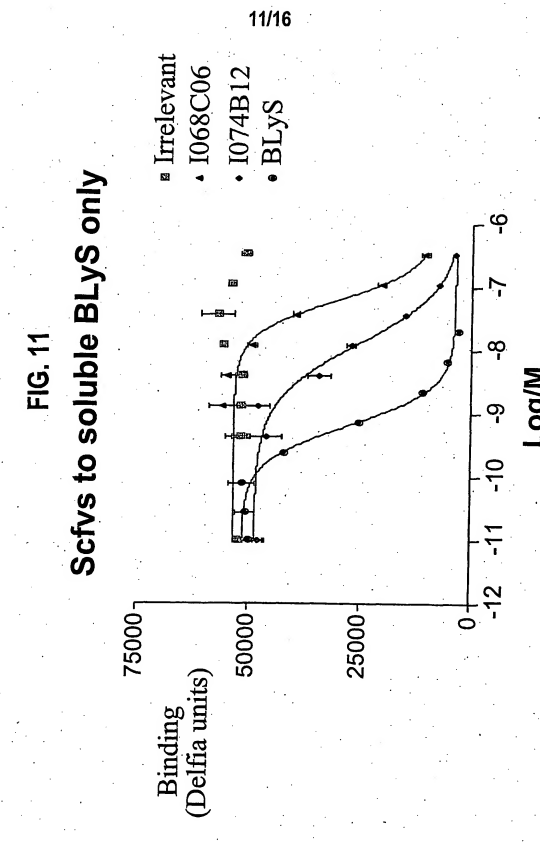




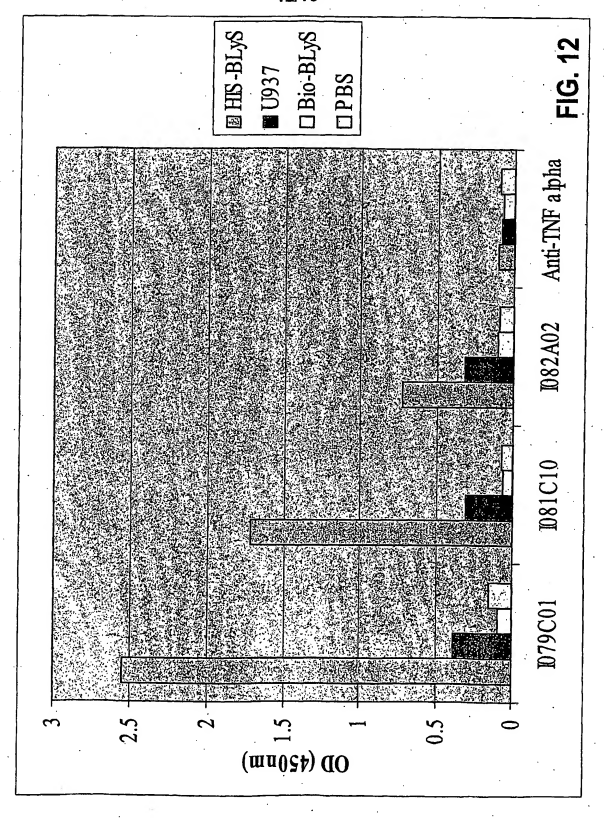




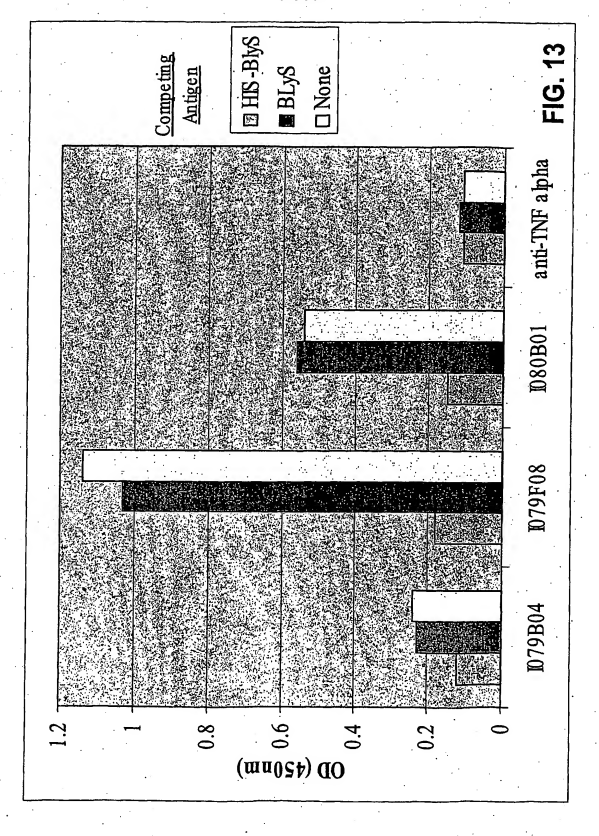




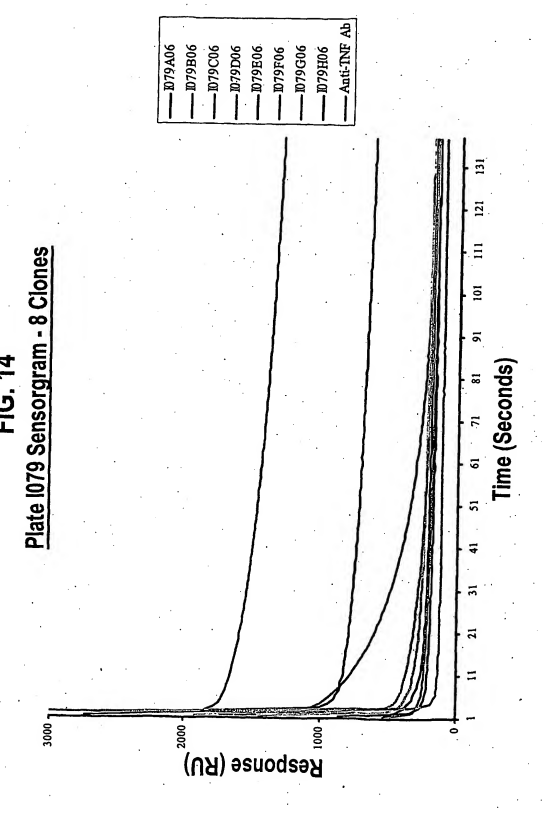
12/16



13/16



14/16



15/16

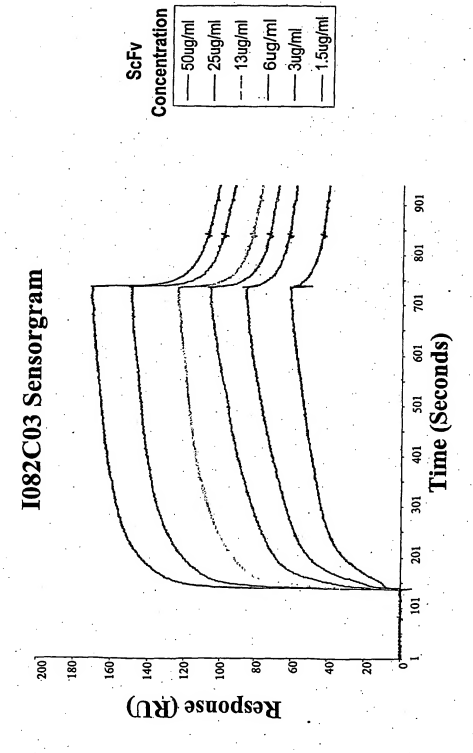
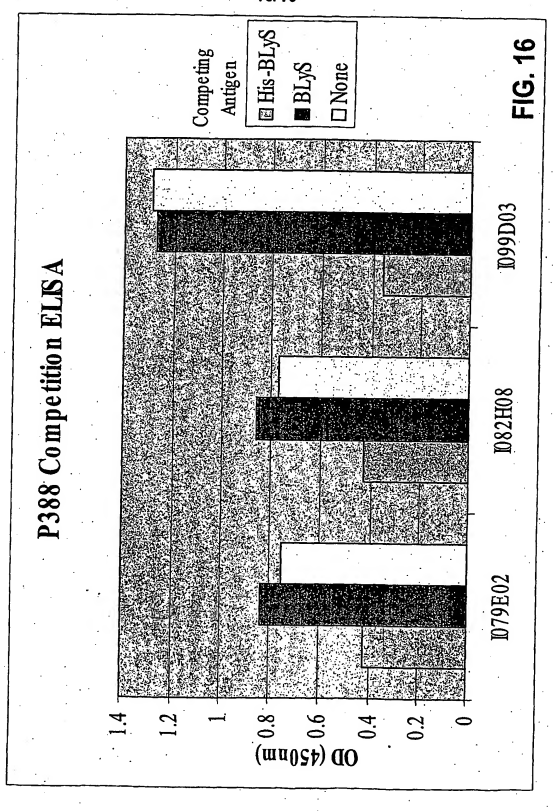


FIG. 15

16/16



SEQUENCE LISTING

- <110> Human Genome Sciences, Inc.
- <120> Antibodies that Immunospecifically Bind BLyS
- <130> PF523PCT
- <140> Not yet assigned
- <141> 2001-06-15
- <150> 60/212,210
- <151> 2000-06-15
- <150> 60/240,816
- <151> 2000-10-17
- <150> 60/276,248
- <151> 2001-03-16
- <150> 60/277,379
- <151> 2001-03-21
- <150> 60/293,499
- <151> 2001-05-25
- <160> 3239
- <170> PatentIn Ver. 2.0
- <210> 1
- <211> 248
- <212> PRT
- <213> Homo sapiens
- <400> 1
- Gln Val Gln Leu Leu Gln Ser Ala Ala Glu Leu Lys Lys Pro Gly Gln
 1 5 10 15
- Ser Leu Lys Ile Ser Cys Lys Gly Ser Gly Phe Thr Phe Thr Tyr
 20 25 30
- Trp Ile Gly Trp Val Arg Gln Leu Pro Gly Lys Gly Leu Glu Trp Met
- Gly Ile Ile Tyr Pro Gly Asp Ser His Thr Thr Tyr Ser Pro Ser Phe
 50 55 60
- Glu Gly His Val Asn Ile Ser Val Asp Lys Ser Ile Asn Thr Ala Tyr
 65 70 75 80

MACIOGEJ

Microsoft Word – Labels4 02/22/06 03:01 PM Joint Discovery Schedule

Realtek's Prelim. Claim Construction Statement -P.L.R. 4-2

Joint Claim Construction and Pre-Hearing Statement - P.L.R. 4-3

3Com's Proposed Terms and Claim Elements P.L.R. 4-1(a)

3Com's Prelim. Claim Construction Statement -P.L.R. 4-2

Draft Joint Claim Construction and Pre-Hearing Statement -P.L.R. 4-3 Realtek's Proposed Terms and Claim Elements P.L.R. 4-1(a)

3Com's Preliminary Claim Construction Statement -P.L.R. 4-2 D-Link's Supp. Prefilminary Claim Construction Statement -P.L.R. 4-2

3Com's Extrinsic Evidence